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A GENE CODING FOR A PROTEIN REGULATING AUREOBASIDIN SENSITIVITY
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[Claim 1] An isolated gene coding for a protein
which regulates aureobasidin sensitivity.

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**ORIGINAL
COMPLETE SPECIFICATION
STANDARD PATENT**

Invention Title: **A GENE CODING FOR A PROTEIN REGULATING
AUREOBASIDIN SENSITIVITY**

The following statement is a full description of this invention, including
the best method of performing it known to us:

GH&CO REF: P17437-C:BJF:RK

[Designation of Document] Specification
[Title of the Invention] A GENE CODING FOR A
PROTEIN REGULATING
AUREOBASIDIN SENSITIVITY

[Detailed Description of the Invention]

[Field of Industrial Application]

This invention relates to a protein regulating the sensitivity to an antimycotic aureobasidin and a gene coding for this protein, namely, a gene coding for a protein regulating aureobasidin sensitivity. The present invention further relates to a series of the uses of the protein and the gene. Furthermore, it relates to an antibody against this protein and the use of the same.

[Prior Art]

Systemic mycoses including candidiasis have increased with an increase in immunocompromised patients in recent years due to, for example, the extended use of immunosuppressive drugs and acquired immunodeficiency syndrome (AIDS), and as opportunistic infection due to microbial substitution caused by the frequent use of wide-spectrum antibacterial antibiotics. Although drugs for treating mycoses such as amphotericin B, flucytosine and azole drugs (for example, fluconazole and miconazole) are now used to cope with this situation, none of them can achieve a satisfactory effect. Also, known diagnostic drugs are insufficient. For candidiasis, in particular, although there have been known several diagnostic drugs (for

example, CAND-TEC for detection of candida antigen and LABOFIT for detection of D-arabinitol), none of them gives any satisfactory results in specificity or sensitivity.

The reasons for the delay in the development of remedies and diagnostic drugs for mycoses as described above are that fungi causing the mycoses are eukaryotic organisms similar to the host (i.e., man) and thus not largely different from man and that knowledges of fungi, in particular, pathogenic fungi are insufficient. Therefore it is difficult to distinguish fungi from man or to selectively kill fungi, which is responsible for the delay in the development of drugs for mycoses.

Recently the application of genetic engineering techniques such as antisense or PCR to the treatment and diagnosis of mycoses has been expected. However known genes which are applicable thereto and/or proteins coded for by these genes are rare (PCT Pamphlet W092/03455). Regarding pathogenic fungi, there have been cloned in recent years an acid protease gene, which has been assumed to participate in the pathogenicity of Candida albicans (hereinafter referred to simply as C. albicans) and Candida tropicalis (hereinafter referred to as C. tropicalis) causing candidiasis [B. Hube et al., J. Med. Vet. Mycol., 29, 129 - 132 (1991); Japanese Patent Laid-Open No. 49476/1993; and G. Togni et al., FEBS Letters, 286, 181 - 185 (1991)], a calmodulin gene of

C. albicans [S.M. Saporito et al., Gene, 106, 43 - 49 (1991)] and a glycolytic pathway enzyme enolase gene of C. albicans [P. Sundstrom et al., J. Bacteriology, 174, 6789 - 6799 (1991)]. However, each of these genes and proteins coded for thereby is either indistinguishable from nonpathogenic fungi and eukaryotic organisms other than fungi or, if distinguishable therefrom, cannot serve as a definite action point for exhibiting any selective toxicity.

Aureobasidin [Japanese Patent Laid-Open No. 138296/1990, No. 22995/1991, No. 220199/1991 and No. 279384/1993, Japanese Patent Application No. 303177/1992, J. Antibiotics, 44 (9), 919 - 924, ibid., 44 (9), 925 - 933, ibid., 44 (11), 1187 - 1198 (1991)] is a cyclic depsipeptide obtained as a fermentation product of a strain Aureobasidium pullulans No. R106. It is completely different in structure from other antimycotics. As Tables 1 and 2 show, aureobasidin A, which is a typical aureobasidin compound, exerts a potent antimycotic activity on various yeasts of the genus Candida including C. albicans which is a pathogenic fungus, Cryptococcus neoformans, Histoplasma capsulatum, Blastomyces dermatitidis and fungi of the genus Aspergillus (Japanese Patent Laid-Open No. 138296/1990) but has an extremely low toxicity in mammal. Thus this compound is expected to be useful as an antimycotic being excellent in selective toxicity.

Hereinafter, Candida, Cryptococcus and
Aspergillus will be abbreviated respectively as C.,
Cr. and A.

[Table 1.]

Test strain	TIMM NO.	MIC(μ g/ml)
<u>C. albicans</u>	0136	≤ 0.04
<u>C. albicans</u> var. <u>stellatoidea</u>	1308	≤ 0.04
<u>C. tropicalis</u>	0312	0.08
<u>C. kefyr</u>	0298	0.16
<u>C. parapsilosis</u>	0287	0.16
<u>C. krusei</u>	0270	≤ 0.04
<u>C. guilliermondii</u>	0257	0.08
<u>C. glabrata</u>	1062	≤ 0.04
<u>Cr. neoformans</u>	0354	0.63
<u>Cr. terreus</u>	0424	0.31
<u>Rhodotorula rubra</u>	0923	0.63
<u>A. fumigatus</u>	0063	20
<u>A. clavatus</u>	0056	0.16

[Table 2]

Test strain	TIMM No.	MIC(μg/ml)
<u>A. nidulans</u>	0112	0.16
<u>A. terreus</u>	0120	5
<u>Penicillium commune</u>	1331	1.25
<u>Trichophyton mentagrophytes</u>	1189	10
<u>Epidemophyton floccosum</u>	0431	2.5
<u>Fonsecaea pedrosoi</u>	0482	0.31
<u>Exophiala werneckii</u>	1334	1.25
<u>Cladosporium bantianum</u>	0343	0.63
<u>Histoplasma capsulatum</u>	0713	0.16
<u>Paracoccidioides brasiliensis</u>	0880	0.31
<u>Geotrichum candidum</u>	0694	0.63
<u>Blastomyces dermatitidis</u>	0126	0.31

[Problems to be Solved by the Invention]

Each of the conventional antimycotics with a weak toxicity shows only a fungistatic effect, which has been regarded as a clinical problem. In contrast, aureobasidin has a fungicidal effect. From this point of view, it has been urgently required to clarify the mechanism of the selective toxicity to fungi of aureobasidin. However this mechanism still remains unknown.

Under these circumstances, the present invention aims at finding a novel protein regulating aureobasidin sensitivity through the clarification of the mechanism of the selective toxicity to fungi of

aureobasidin. Accordingly, the present invention aims at finding a gene coding for a protein regulating aureobasidin sensitivity, providing a process for cloning this gene and the protein regulating aureobasidin sensitivity which is encoded by this gene, further providing an antisense DNA and an antisense RNA of this gene, providing a nucleic acid probe being hybridizable with this gene, providing a process for detecting this gene with the use of the nucleic acid probe, providing a process for producing the protein regulating aureobasidin sensitivity by using this gene and providing an antibody against the protein regulating aureobasidin sensitivity, and a process for detecting the protein regulating aureobasidin sensitivity by using this antibody.

[Means for Solving the Problems]

The present invention may be summarized as follows. Namely, the first invention of the present invention relates to an isolated gene coding for a protein regulating aureobasidin sensitivity, that is, a gene regulating aureobasidin sensitivity. The second invention relates to a process for cloning a gene regulating aureobasidin sensitivity which is characterized by using the gene regulating aureobasidin sensitivity of the first invention or a part thereof as a probe. The third invention relates to a nucleic acid probe which is hybridizable with a gene regulating aureobasidin sensitivity and comprises a sequence consisting of 15 or more bases. The fourth

invention relates to an antisense DNA of a gene regulating aureobasidin sensitivity. The fifth invention relates to an antisense RNA of a gene regulating aureobasidin sensitivity. The sixth invention relates to a recombinant plasmid having a gene regulating aureobasidin sensitivity contained therein. The seventh invention relates to a transformant having the above-mentioned plasmid introduced thereto. The eighth invention relates to a process for producing a protein regulating aureobasidin sensitivity by using the above-mentioned transformant. The ninth invention relates to an isolated protein regulating aureobasidin sensitivity. The tenth invention relates to an antibody against a protein regulating aureobasidin sensitivity. The eleventh invention relates to a process for detecting a protein regulating aureobasidin sensitivity by using the above-mentioned antibody. The twelfth invention relates to a process for detecting a gene regulating aureobasidin sensitivity by the hybridization which is characterized by using the nucleic acid probe of the third invention of the present invention. The thirteenth invention relates to a process for screening an antimycotic by using the above-mentioned transformant or a protein regulating aureobasidin sensitivity.

The present inventors have found out that fungi such as Schizosaccharomyces pombe (hereinafter referred to simply as Schizo. pombe) and Saccharomyces

cerevisiae (hereinafter referred to simply as S. cerevisiae) and, further, mammalian cells such as mouse lymphoma EL-4 cells are sensitive to aureobasidin, as Table 3 shows.

[Table 3]

Test strain or cell	MIC(μg/ml)
<u>Schizo. pombe</u>	0.08
<u>S. cerevisiae</u>	0.31
mouse lymphoma EL-4	10
mouse lymphoma L5178Y	100
NRK-49F	12.5

The present inventors have mutagenized a wild-type strain of Schizo. pombe or S. cerevisiae, sensitive to aureobasidin, to thereby give resistant mutants. We have further successfully isolated a gene capable of conferring aureobasidin resistance (a resistant gene) from these resistant mutants and another gene capable of imparting aureobasidin sensitivity (a sensitive gene) from the corresponding sensitive cells.

Furthermore, we have disclosed the existence of a protein encoded by each of these genes. By culturing cells which have been transformed by introducing the above-mentioned gene, we have succeeded in the expression of this gene. Furthermore, we have successfully found out a novel gene regulating aureobasidin sensitivity from another fungus being

sensitive to aureobasidin by using a DNA fragment of the above-mentioned gene as a probe. In addition, we have clarified that the gene regulating aureobasidin sensitivity is essentially required for the growth of the cells and found out that the detection of this gene or a protein which is a gene product thereof with an antibody enables the diagnosis of diseases caused by these cells, for example, mycoses induced by fungi, and that an antisense DNA or an antisense RNA, which inhibits the expression of the gene regulating aureobasidin sensitivity being characteristic to the cells, is usable as a remedy for diseases caused by these cells, for example, mycoses induced by fungi, thus completing the present invention.

That is to say, pathogenic fungi listed in Tables 1 and 2 and fungi and mammalian cells listed in Table 3, each having a sensitivity to aureobasidin, each carries a protein regulating aureobasidin sensitivity and a gene coding for this protein. The term "a protein regulating aureobasidin sensitivity" as used herein means a protein which is contained in an organism having a sensitivity to aureobasidin. This protein is required for the expression of the sensitivity or resistance to aureobasidin. As a matter of course, a protein having 35% or more homology with the above-mentioned protein and having a similar function is also a member of the protein regulating aureobasidin sensitivity according to the present invention. Furthermore, proteins obtained by

modifying these proteins by the genetic engineering procedure are members of the protein regulating aureobasidin sensitivity according to the present invention. A gene regulating aureobasidin sensitivity means a gene which codes for such a protein regulating aureobasidin sensitivity as those described above and involves both of sensitive and resistant genes.

The first invention of the present invention relates to a gene regulating aureobasidin sensitivity. This gene can be isolated in the following manner.

First, aureobasidin sensitive cells (a wild-type strain) is mutagenized to thereby induce a resistant strain. From chromosome DNA or cDNA of this resistant strain, a DNA library is prepared and a gene capable of conferring a resistance (a resistant gene) is cloned from this library. Then a DNA library of a wild strain is prepared and a DNA molecule being hybridizable with the resistant gene is isolated from this library and cloned. Thus a sensitive gene can be isolated.

The mutagenesis is performed by, for example, treating with a chemical such as ethylmethane sulfonate (EMS) or N-methyl-N'-nitro-N-nitrosoguanidine (MNNG) or by ultraviolet or other radiation. The cell that has acquired the resistance can be screened by culturing the mutagenized cells in a nutritional medium containing aureobasidin at an appropriate concentration under appropriate conditions. The resistant strain thus

obtained may vary depending on the method and conditions selected for the mutagenesis. Also, strains differing in the extent of resistance from each other can be separated by changing the aureobasidin concentration or a temperature-sensitive resistant strain can be isolated by changing the temperature in the step of screening. There are a number of mechanisms of resistance to aureobasidin. Accordingly, a number of resistant genes can be isolated by genetically classifying these various resistant strains. In the case of a yeast, the classification may be performed by the complementation test. Namely, resistant strains are prepared from haploid cells. Next, diploid cells can be obtained by crossing resistant strains differing in mating type from each other. Then spores formed from these diploids are examined by the tetrad analysis.

As typical examples of the genes regulating aureobasidin sensitivity (named aur) according to the present invention, aur1 and aur2 genes may be cited. Typical examples of the aur1 gene include spa^r1 gene isolated from Schizo. pombe and sca^r1 gene isolated from S. cerevisiae, while typical examples of the aur2 gene include sca^r2 gene isolated from S. cerevisiae.

Now, resistant genes (spa^r1^r, sca^r1^r and sca^r2^r) isolated from resistant mutants by the present inventors and sensitive genes (spa^r1^s, sca^r1^s and sca^r2^s) isolated from sensitive wild-type strains will be described.

Fig. 1 shows a restriction enzyme map of the genes spaurl^R and spaurl^S regulating aureobasidin sensitivity, Fig. 2 shows a restriction enzyme map of scaur1^R and scaur1^S and Fig. 3 shows a restriction enzyme map of scaur2^R and scaur2^S.

Schizo. pombe, which is sensitive to aureobasidin, is mutagenized with EMS and a genomic library of the resistant strain thus obtained is prepared. From this library, a DNA fragment containing a resistant gene (spaurl^R) and having the restriction enzyme map of Fig. 1 is isolated. This gene has a nucleotide sequence represented by SEQ ID No. 1 in Sequence Listing. The amino acid sequence of a protein encoded by this gene, which is estimated on the basis of this nucleotide sequence, is the one represented by SEQ ID No. 2 in Sequence Listing. By the hybridization with the use of this resistant gene as a probe, a DNA fragment containing a sensitive gene (spaurl^S) and having the restriction enzyme map of Fig. 1 is isolated from a sensitive strain. This gene has a nucleotide sequence represented by SEQ ID No. 3 in Sequence Listing. The amino acid sequence of a protein encoded by this gene, which is estimated on the basis of this nucleotide sequence, is the one represented by SEQ ID No. 4 in Sequence Listing. A comparison between the sequences of SEQ ID No. 3 and SEQ ID No. 1 reveals that a mutation from G to T occurs at the base at the position 1053, while a comparison between the sequences of SEQ ID No. 4 and SEQ ID No. 2 reveals

that glycine at the residue 240 is converted into cysteine at the amino acid level, thus giving rise to the resistance.

Also, S. cerevisiae, which is sensitive to aureobasidin, is mutagenized with EMS and genomic libraries of two resistant strains thus obtained are prepared. From one of these libraries, a DNA fragment containing a resistant gene (*scaur1^R*) as a dominant mutant and having the restriction enzyme map of Fig. 2 is isolated, while a DNA fragment containing a resistant gene (*scaur2^R*) and having the restriction enzyme map of Fig. 3 is isolated from another library.

The nucleotide sequence of the coding region for the protein of the *scaur1^R* gene is the one represented by SEQ ID No. 5 in Sequence Listing. The amino acid sequence of the protein encoded by this gene, which is estimated on the basis of the above nucleotide sequence, is the one represented by SEQ ID No. 6 in Sequence Listing. By the hybridization with the use of this resistant gene *scaur1^R* as a probe, a DNA fragment containing a sensitive gene (*scaur1^S*) and having the restriction enzyme map of Fig. 2 is isolated from a sensitive strain. This gene has a nucleotide sequence represented by SEQ ID No. 7 in Sequence Listing. The amino acid sequence of a protein encoded by this gene, which is estimated on the basis of this nucleotide sequence, is the one represented by SEQ ID No. 8 in Sequence Listing. A comparison between the sequences of SEQ ID No. 7 and SEQ ID No. 5 reveals that a

mutation from T to A occurs at the base at the position 852, while a comparison between the sequences of SEQ ID No. 8 and SEQ ID No. 6 reveals that phenylalanine at the residue 158 is converted into tyrosine at the amino acid level, thus giving rise to the resistance. The spaurl gene has a 58% homology with the scaurl gene at the amino acid level. Thus it is obvious that they are genes coding for proteins having similar functions to each other. When genes and proteins being homologous in sequence with the spaurl and scaurl genes and with the proteins encoded thereby are searched from a data base, none having a homology of 35% or above is detected. Accordingly, it is clear that these genes and the proteins encoded thereby are novel molecules which have never been known hitherto.

By the hybridization with the use of the DNA fragment of the resistant gene scaur2^R as a probe, a DNA fragment containing a sensitive gene (scaur2^S) and having the restriction enzyme map of Fig. 3 is isolated from a sensitive strain.

The nucleotide sequence of this sensitive gene is the one represented by SEQ ID No. 9 in Sequence Listing and the amino acid sequence of the protein encoded by this gene, which is estimated on the basis of this nucleotide sequence, is the one represented by SEQ ID No. 10 in Sequence Listing. As the result of the homology search with the scaur2^S gene and the protein encoded thereby, it has been found out that cystic fibrosis transmembrane conductance regulator

(CFTR) of mammals alone has a homology as low as 31%.

Compared with this CFTR, however, the part having a high homology is limited to the region around the domain of the nucleotide binding. It is therefore obvious that the protein encoded by the scaur2^s gene is a protein which is completely different from CFTR in function and has never been known hitherto.

In order to prove the importance of the aurl gene in the growth of cells, genes for disrupting the aurl as shown in Fig. 4 and Fig. 5, in which genes coding for orotidine-5'-phosphate decarboxylase (*ura4*⁺ in the case of Schizo. pombe, while *URA3* in the case of S. cerevisiae) have been introduced midway in the aurl gene, are prepared. When these aurl disrupted genes are introduced into Schizo. pombe and S. cerevisiae respectively, the cells having the aurl disrupted genes cannot grow at all. Thus it has been revealed that these genes and the proteins encoded thereby are essentially required for the growth of the yeast cells.

As the above examples clearly show, a gene regulating aureobasidin sensitivity can be isolated by using a organism having sensitivity to aureobasidin as a starting material and by carrying out the cloning with the use of various mutagenesis methods and/or screening methods depending on the organisms or the methods. Also, genes being hybridizable with the above-mentioned genes are involved in the scope of the first invention of the present invention. A gene

regulating aureobasidin sensitivity can be isolated by the following method. The genomic DNA library of an organism having sensitivity to aureobasidin is integrated into, for example, a high-expression vector of a yeast and transformed into the yeast. Then a clone having aureobasidin resistance is selected from the transformants and DNA is recovered from this clone. Thus the resistant gene can be obtained. As a matter of course, genes obtained by modifying some part of the gene regulating aureobasidin sensitivity thus obtained by some chemical or physical methods are involved in the scope of the first invention of the present invention.

The second invention of the present invention relates to a process for cloning a gene regulating aureobasidin sensitivity which is characterized by using the gene regulating aureobasidin sensitivity of the first invention of the present invention or a part thereof as a probe. Namely, by screening by the hybridization method or the polymerase chain reaction (PCR) method with the use of a part (consisting of at least 15 oligonucleotides) or the whole of the gene as obtained above, a gene coding for a protein having a similar function can be isolated.

For example, a pair of primers of SEQ ID No. 11 and SEQ ID No. 12 in Sequence Listing are synthesized on the basis of the DNA nucleotide sequence of the *spaur1^R* gene represented by SEQ ID No. 1. Then PCR is performed by using cDNA of C. albicans, which is a

pathogenic fungus, as a template with the use of the above-mentioned primers. The PCR is carried out and the PCR products are electrophoresed on an agarose gel and stained with ethidium bromide. In Fig. 6, the lanes 1, 2 and 3 show the results obtained by using cDNA of C. albicans, cDNA of S. cerevisiae and cDNA of Schizo. pombe as a template, respectively. As shown in Fig. 6, a certain DNA fragment is specifically amplified.

By screening the genomic DNA library of C. albicans with the use of this DNA fragment as a probe, a DNA molecule having a gene (caaurl), which has the same function as that of the spaurl and scaurl genes and having the restriction enzyme map of Fig. 7 is obtained. The nucleotide sequence of this caaurl gene is the one represented by SEQ ID No. 13 in Sequence Listing and the amino acid sequence of the protein encoded by this gene, which has been estimated on the basis of the above nucleotide sequence, is the one represented by SEQ ID No. 14 in Sequence Listing. It has a high homology with the proteins encoded by the spaurl and scaurl genes.

By screening the genomic DNA library of C. albicans with the use of a DNA fragment comprising the whole length or a part of the scaur2^s gene represented by SEQ ID No. 9 in Sequence Listing as a probe, a DNA fragment containing gene (caaurl2), which has the same function as that of the scaur2 gene, and having the restriction enzyme map of Fig. 8 is obtained. The

nucleotide sequence of a part of this caaur2 gene is the one represented by SEQ ID No. 15 in Sequence Listing and the amino acid sequence of the region encoded by this gene, which has been estimated on the basis of this nucleotide sequence, is the one represented by SEQ ID No. 16 in Sequence Listing. It has a high homology with the corresponding region of the protein encoded by the scaur2 gene.

The third invention of the present invention relates to an oligonucleotide comprising 15 or more bases which serves as the above-mentioned nucleic acid probe and is hybridizable with the gene regulating aureobasidin sensitivity, for example, the DNA fragment having the restriction enzyme map as shown in Fig. 1, Fig. 2 or Fig. 3. This nucleic acid probe is usable in, for example, the hybridization in situ, the identification of a tissue wherein the above-mentioned gene can be expressed, and the confirmation of the presence of a gene or mRNA in various vital tissues. This nucleic acid probe can be prepared by ligating the above-mentioned gene or a gene fragment to an appropriate vector, introducing it into a bacterium, allowing it to replicate in the bacterium, extracting from a disrupted cell suspension, cleaving with a restriction enzyme capable of recognizing the vector-ligating site, electrophoresing and then excising from the gel. Alternatively, this nucleic acid probe can be constructed by the chemical synthesis with the use of a DNA synthesizer or gene amplification techniques by

PCR on the basis of the nucleotide sequence of SEQ ID.

Nos. 1, 3, 5, 7, 9, 13, 15 or 21 in Sequence Listing.

This nucleic acid probe can be labeled with a radioisotope or a fluorescent substance to thereby elevate the detection sensitivity at the use.

The fourth invention of the present invention relates to an antisense DNA of the above-mentioned gene regulating aureobasidin sensitivity, while the fifth invention of the present invention relates to an antisense RNA thereof. The introduction of the antisense DNA or antisense RNA into cells makes it possible to control the expression of the gene regulating aureobasidin sensitivity.

As examples of the antisense DNA to be introduced, antisense DNAs corresponding to the genes regulating aureobasidin sensitivity of SEQ ID Nos. 1, 3, 5, 7, 9, 13, 15 or 21 in Sequence Listing and some parts thereof may be cited. SEQ ID No. 17 in Sequence Listing shows an example of this antisense DNA. It represents the sequence of an antisense DNA of the gene regulating aureobasidin sensitivity of SEQ ID No. 1 in Sequence Listing. A fragment obtained by appropriately cleaving some part of such an antisense DNA, and a DNA synthesized depending on such an antisense DNA sequence may be used as the antisense DNA.

As examples of the antisense RNA to be introduced, antisense RNAs corresponding to the genes regulating aureobasidin sensitivity of SEQ ID Nos. 1, 3, 5, 7, 9,

13, 15 or 21 in Sequence Listing and some parts thereof may be cited. SEQ ID No. 18 in Sequence Listing shows an example of this antisense RNA. It represents the sequence of an antisense RNA of the gene regulating aureobasidin sensitivity of SEQ ID NO. 1 in Sequence Listing. A fragment obtained by appropriately cleaving some part of such an antisense RNA, an RNA synthesized depending on such an antisense RNA sequence, and an RNA prepared with RNA polymerase in an in vitro transcription system by using the DNA corresponding to the gene regulating aureobasidin sensitivity of SEQ ID NO. 1 or SEQ ID NO. 3 in Sequence Listing or a part thereof may be used as the antisense RNA.

These antisense DNA and antisense RNA may be chemically modified so as to prevent degradation in vivo or to facilitate passage through a cell membrane. A substance capable of inactivating mRNA, for example, ribozyme may be linked thereto. The antisense DNA and antisense RNA thus prepared are usable in the treatment of various diseases such as mycoses accompanied by an increase in the amount of mRNA coding for a protein regulating aureobasidin sensitivity.

The sixth invention of the present invention relates to a recombinant plasmid having a gene coding for a protein regulating aureobasidin sensitivity being integrated into an appropriate vector. For example, a plasmid, in which a gene regulating

aureobasidin sensitivity gene has been integrated into an appropriate yeast vector, is highly useful as a selection marker gene, since a transformant can be easily selected thereby with the guidance of the chemical resistance by using *aureobasidin*.

Also, the recombinant plasmid can be stably carried by, for example, Escherichia coli. Examples of vectors which are usable in this case include pUC118, pWH5, pAU-PS, Traplex119 and pTV118. pAU-PS having the *spaur1^s* gene integrated therein is named pSPAR1. pWH5 having the *spaur1^s* gene integrated therein is named pSCAR1. pWH5 having the *scaur2^R* gene integrated therein is named pSCAR1. Traplex119 vector having the *caauml* gene integrated therein is named pCAAR1. pTV118 vector having a part of the *caauml2* gene integrated therein is named pCAAR2N. Each of these recombinant plasmids is transformed into E. coli. It is also possible to express these plasmids in an appropriate host. Such a gene is reduced exclusively into the open reading frame (ORF) to be translated into a protein by cleaving with an appropriate restriction enzyme, if necessary, and then bound to an appropriate vector. Thus an expression recombinant plasmid can be obtained. When E. coli is used as the host, plasmids such as pTV118 may be used as a vector for the expression plasmid. When a yeast is used as the host, plasmids such as pYES2 may be used as the vector. When mammalian cells are used as the host, plasmids such as pMAMneo may be used as the vector.

The seventh invention of the present invention relates to a transformant having the above-mentioned recombinant plasmid which has been introduced into an appropriate host. As the host, E. coli, yeasts and mammalian cells are usable. E. coli JM109 transformed by pSPAR1 having the spaurl^s gene integrated therein has been named and designated as Escherichia coli JM109/pSPAR1 and deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology (1-3, Higashi 1 chome Tsukuba-shi Ibaraki-ken 305, JAPAN) on 13 April 1993, in accordance with the Budapest Treaty under the accession number FERM BP-4485. E. coli HB101 transformed by pSCAR1 having the scaurl^s gene integrated therein has been named and designated as Escherichia coli HB101/pSCAR1 and deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on 13 April 1993 in accordance with the Budapest Treaty under the accession number FERM BP-4483. E. coli HB101 transformed by pSCAR2 having the scaur2^s gene integrated therein has been named and designated as Escherichia coli HB101/pSCAR2 and deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on 13 April 1993 in accordance with the Budapest Treaty under the accession number FERM BP-4484. E. coli HB101 transformed by pCCAR1 having the caaurl^s gene integrated therein has been named and designated as Escherichia coli HB101/pCCAR1 and deposited at

National Institute of Bioscience and Human-Technology,
Agency of Industrial Science and Technology on 1
December 1993 in accordance with the Budapest Treaty
under the accession number FERM BP-4482. E. coli HB101
transformed by pCAAR2N having a part of the caaur2
gene integrated therein has been named and designated
as Escherichia coli HB101/pCAAR2N and deposited at
National Institute of Bioscience and Human-Technology,
Agency of Industrial Science and Technology on 1
December 1993 in accordance with the Budapest Treaty
under the accession number FERM BP-4481.

A transformant capable of expressing a protein
regulating aureobasidin sensitivity can be obtained by
transforming a expression recombinant plasmid into an
appropriate host, as described above. For example, a
yeast having a recombinant plasmid as shown in Fig. 9
introduced thereinto is usable for this purpose.

The eighth invention of the present invention
relates to a process for producing a protein
regulating aureobasidin sensitivity which comprises
incubating a transformant according to the sixth
invention of the present invention, which contains a
gene coding for this protein, in an appropriate
nutritional medium, allowing the expression of the
protein, then recovering the protein from the cells or
the medium and purifying the same. For the expression
of the gene coding for this protein, E. coli, a yeast
or mammalian cells are employed as a host. When the
yeast having the recombinant plasmid of Fig. 9 is

incubated in a medium containing galactose, for example, the protein regulating aureobasidin sensitivity which is encoded by the *scaur1^s* gene can be expressed.

The ninth invention of the present invention relates to an isolated protein regulating aureobasidin sensitivity. As examples of such a protein, those encoded by the above-mentioned *spaur1*, *scaur1*, *scaur2*, *caaurl* and *caaurl2* genes can be cited.

The *spaur1^s* gene codes for a protein having an amino acid sequence represented by SEQ ID No. 4 in Sequence Listing, while the *scaur1^s* gene codes for a protein having an amino acid sequence represented by SEQ ID No. 8 in Sequence Listing. By the northern hybridization with the use of a DNA fragment of the *spaur1* gene as a probe, mRNAs are detected from a sensitive strain (Fig. 10). Thus the expression of the *spaur1* gene is confirmed.

Fig. 10 is an autoradiogram showing the results of the northern hybridization wherein mRNAs obtained from cells of a sensitive strain of Schizo. pombe in the logarithmic growth phase (lane 1), cells of a resistant strain in the logarithmic growth phase (lane 2), cells of the sensitive strain in the stationary phase (lane 3) and cells of the resistant strain in the stationary phase (lane 4) are electrophoresed on a 1.2% agarose gel containing formaldehyde.

The tenth invention of the present invention relates to an antibody against the above-mentioned

protein regulating aureobasidin sensitivity. For example proteins having amino acid sequences of SEQ ID Nos. 2, 4, 6, 8, 10, 14, 16 or 22 in Sequence Listing and peptides comprising some parts of these amino acid sequences may be used as an antigen. The former antigens can be prepared through the expression in a transformant followed by purification, while the latter antigens can be synthesized on, for example, a marketed synthesizer. The antibody is produced by the conventional method. For example, an animal such as a rabbit is immunized with the above-mentioned protein or a peptide fragment together with an adjuvant to thereby give a polyclonal antibody. A monoclonal antibody can be produced by fusing antibody-producing B cells, which have been obtained by immunizing with an antigen, with myeloma cells, screening hybridomas producing the target antibody, and incubating these cells. As will be described hereinafter, these antibodies are usable in the treatment and diagnosis for animal and human diseases in which the above-mentioned proteins participate, such as mycoses.

For example, a peptide corresponding to the part of the 103- to 113-positions in the amino acid sequence of SEQ ID No. 8 is synthesized on a synthesizer and then bound to a carrier protein. Then a rabbit is immunized therewith and thus a polyclonal antibody is obtained. In the present invention, keyhole limpet hemocyanin (KLH) is used as the carrier protein. Alternatively, bovine serum albumin and

ovalbumin are usable therefor.

The eleventh invention of the present invention relates to a process for detecting a protein regulating aureobasidin sensitivity by using the above-mentioned antibody. The detection can be carried out by detecting the binding of the antibody to the protein or measuring the amount of binding. For example, the protein or the cells producing the same can be detected by treating with a fluorescence-labeled antibody and then observing under a fluorescence microscope. The amount of the antibody bound to the protein can be measured by various known methods. For example, S. cerevisiae cells are stained by the immunofluorescent antibody technique by using the above-mentioned antibody and a secondary antibody such as FITC-labeled antirabbit antibody. Thus it is clarified that the protein encoded by the *scaur1* gene is distributed all over the cells. Further, a yeast having the recombinant plasmid of Fig. 9 introduced thereto is incubated in a medium containing glucose or galactose. The cells thus obtained are disrupted with glass beads and proteins are solubilized. Then these proteins are separated by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and the western blotting is carried out in the conventional manner by using the above-mentioned polyclonal antibody and peroxidase-labeled anti-rabbit antibody. Consequently, the protein encoded by the *scaur1* gene can be detected, as Fig. 11 shows.

Fig. 11 shows the results of the western blotting wherein the proteins prepared from the cells obtained by the incubation in the presence of glucose (lane 1) or galactose (lane 2) are subjected to SDS-PAGE. A main band binding to the polyclonal antibody of the present invention is detected at around 38 kDa.

The twelfth invention of the present invention relates to a process for detecting a gene regulating aureobasidin sensitivity, for example, mRNA at the expression of a protein, by using the above-mentioned oligonucleotide as a nucleic acid probe. This process is applicable to the diagnosis for various diseases, including mycoses, associated with an abnormal amount of mRNA coding for the protein. For example, nucleic acids are precipitated from disrupted cells and mRNA is hybridized with a radioisotope-labeled nucleic acid probe on a nitrocellulose membrane. The amount of binding can be measured by autoradiography (Fig. 10) or with a scintillation counter.

The thirteenth invention of the present invention relates to a process for efficient screening of a novel antimycotic by using the transformant of the seventh invention of the present invention or the protein regulating aureobasidin sensitivity of the ninth invention of the present invention. For example, a drug exerting its effect on the protein or the gene of the present invention can be efficiently found out through a comparison of the activity on a transformant

containing a sensitive gene with the activity on a transformant containing a resistant gene or a comparison between the activities on transformants differing in expression level from each other. Also, the screening can be efficiently carried out by measuring the affinity for the protein of the present invention, for example, the activity of inhibiting the binding of radiolabeled-aureobasidin to the protein.

[Brief Description of the Drawings]

[Fig. 1]

Restriction enzyme map of the genes $spaurl^R$ and $spaurl^S$ regulating aureobasidin sensitivity.

[Fig. 2]

Restriction enzyme map of $scaur1^R$ and $scaur1^S$.

[Fig. 3]

Restriction enzyme map of $scaur2^R$ and $scaur2^S$.

[Fig. 4]

Structure of a DNA for disrupting the Schizo. pombe $spaurl^S$ gene.

[Fig. 5]

Structure of a DNA for disrupting the S. cerevisiae $scaur1^S$ gene.

[Fig. 6]

Results of the detection of the $aurl$ gene $caaurl$ carried by C. albicans by the PCR method.

[Fig. 7]

Restriction enzyme map of the $caaurl$ gene carried by C. albicans.

[Fig. 8]

Restriction enzyme map of the caaur2 gene.

[Fig. 9]

Structure of a plasmid YEpSCARW3 for expressing the scaur1 gene.

[Fig. 10]

Results of the northern hybridization of the scaur1 gene of Schizo. pombe.

[Fig. 11]

Results of the detection of the scaur1 protein by using an antibody.

[Fig. 12]

Restriction enzyme map of pAR25.

[Examples]

To further illustrate the present invention in greater detail, the following Examples will be given. However it is to be understood that the present invention is not restricted thereto.

Example 1: Cloning of a gene regulating aureobasidin sensitivity originating in fission yeast

Schizo. pombe

I-a) Separation of aureobasidin-resistant mutant of Schizo. pombe

About 1×10^8 cells of a Schizo. pombe haploid cell strain JY745 (mating type h, genotype ade6-M210, leul, ura4-D18) exhibiting a sensitivity to aureobasidin at a concentration of 0.08 µg/ml were suspended in 1 ml of a phosphate buffer containing

0.9% NaCl. Then the cells were mutagenized with EMS at a final concentration of 3% at 30°C for 90 minutes. After neutralizing by adding 8 ml of 5% sodium thiosulfate, the cells thus treated were harvested by centrifugation (2500 r.p.m., 5 minutes), washed twice with 6 ml of physiological saline and then suspended in 2 ml of a YEL medium (3% of glucose, 0.5% of yeast extract). The suspension was incubated at 30°C for 5 hours under stirring and then spreaded on a YEA plate (the YEL medium containing 1.5% of agar) containing 5 µg/ml of aureobasidin A. After incubating at 30°C for 3 to 4 days, two or three aureobasidin-resistant colonies were formed per 1×10^8 cells. After carrying out the mutagenesis several times, five clone mutants, i.e., THR01, THR04, THR05, THR06 and THR07 were obtained. These mutants were resistant to more than 25 µg/ml of aureobasidin A but the same as the parent strain in the sensitivity to cycloheximide and amphotericin B. Therefore it is estimated that they are not mutants having a multiple drug resistance but ones having a resistance specific to aureobasidin.

1-b) Genetic analysis

Each of the above-mentioned resistant strains THR01, THR04, THR05, THR06 and THR07 was crossed with normal cells of Schizo. pombe LH121 strain (mating type h, genotype ade6-M216, ura4-D18) differing in mating type. Diploid cells obtained were examined about the resistance to aureobasidin. Similar to the

resistant strains, the five diploids formed by crossing the resistant strains with the normal one were resistant to 25 µg/ml of aureobasidin A, thus proving that these resistant mutations were dominant.

To perform the tetrad analysis, the above-mentioned diploids were subsequently inoculated on an MEA medium (3% of malt extract, 2.5% of agar) for sporulation and incubated at 25°C for 2 days. Prior to the meiosis, the diploid cells replicated DNA on the MEA medium and then underwent the meiosis to form asci each containing four ascospores of the haploid. These spores were separated with a micromanipulator and allowed to germinate on the YEA plate, followed by the formation of colonies. Then the resistance to aureobasidin of these colonies was examined. Among four spores contained in an ascus, the separation of the sensitivity versus the resistance showed 2 : 2. This result indicates that the aureobasidin resistant mutation was induced by a mutation in single gene.

Further, the complementation test was performed in order to confirm whether the resistant genes of the above-mentioned five mutants were identical with each other or not. For example, a mutant of the mating type "h", which had been obtained by crossing the mutant THR01 with the LH121 strain in the above tetrad analysis, was crossed with another variant THR04 (mating type "h") on the MEA plate as described above and, after sporulation, the tetrad analysis was carried out. As a result, all of the colonies formed

from four ascospores showed resistance to aureobasidin, which indicates that the mutational genes of THR01 and THR04 are the same with each other. Similarly, the five mutants were examined and it was thus found out that all mutations occurred on the same gene. This gene regulating aureobasidin sensitivity is named spaurl, the normal gene (sensitive gene) is named spaurl^s and the mutational gene (resistant gene) is named spaurl^r.

1-c) Preparation of genomic library of aureobasidin resistant strain

Genomic DNA was extracted and purified from the aureobasidin resistant strain THR01 by the method of P. Philippson et al. [Methods in Enzymology, 194, 169 - 175 (1991)]. The purified genomic DNA (8 µg) was partially digested by treating with 5 U of a restriction enzyme HindIII at 37°C for 10 minutes, deproteinized with phenol/chloroform and precipitated with ethanol. The partially digested DNA was electrophoresed on a 0.8% agarose gel and DNA in the region of 3 to 15 kb was extracted. The DNA thus obtained was ligated with a yeast-E. coli shuttle vector pAU-PS (2 µg) which had been completely digested with HindIII by using a DNA ligation kit (manufactured by Takara Shuzo Co., Ltd.) and then transformed into E. coli HB101. Thus a genomic library of the aureobasidin resistant strain was formed. E. coli containing this genomic library was incubated in 50 ml of an LB medium (1% of bacto tryptone, 0.5% of

bacto yeast extract, 0.5% of sodium chloride) containing 100 µg/ml of ampicillin and 25 µg/ml of tetracycline at 37°C overnight. Then a plasmid was recovered and purified from the E. coli cells.

1-d) Expression and cloning of aureobasidin resistant gene *spaur1^R*

The plasmid originating in the genomic library of the aureobasidin resistant strain as prepared above was transformed into a strain Schizo. pombe JY745 by the method of Okazaki et al. [Nucleic Acid Research 18, 6485 - 6489 (1990)]. The transformed cells were spreaded on a minimum medium SD plate [0.67% of yeast nitrogen base without amino acids (manufactured by Difco), 2% of glucose, 2% of agar] containing 75 µg/ml of adenine sulfate and 50 µg/ml of leucine. After incubating at 30°C for 3 to 4 days, the colonies thus formed were replicated onto an SD plate containing 5 µg/ml of aureobasidin A, 75 µg/ml of adenine sulfate and 50 µg/ml of leucine. It is conceivably that a colony propagated on this plate may have the plasmid containing the aureobasidin resistant gene. This colony was inoculated into 5 ml of a liquid SD medium containing 75 µg/ml of adenine sulfate and 50 µg/ml of leucine. After incubating at 30°C for 2 days, the plasmid was recovered from the propagated cells by the method of I. Hagan et al. [J. Cell Sci., 91, 587 - 595 (1988)]. Namely, the cells were harvested from the culture (5 ml) by centrifugation and then suspended in 1.5 ml of 50 mM citrate/phosphate buffer containing

1.2 M of sorbitol and 2 mg/ml of Zymolyase. Then the suspension was maintained at 37 °C for 60 minutes. The cells were collected by centrifuging at 3,000 r.p.m. for 30 seconds and suspended in 300 µl of a TE [10 mM of Tris-HCl, pH 8, 1 mM of EDTA] solution. After adding 35 µl of 10% SDS, the mixture was maintained at 65°C for 5 minutes. After adding 100 µl of 5 M potassium acetate, the mixture was allowed to stand in ice for 30 minutes. Then it was centrifuged at 10,000 r.p.m. at 4°C for 10 minutes and a plasmid DNA was purified from the supernatant by using EASYTRAP™ (manufactured by Takara Shuzo Co., Ltd.).

This plasmid was transformed into E. coli HB101 and a plasmid DNA was prepared from E. coli colonies formed on an LB medium containing ampicillin and tetracycline. This plasmid, which contained a DNA of 4.5 kb, was named pAR25. Fig. 12 shows the restriction enzyme map of the DNA of 4.5 kb in pAR25. To specify the gene region, HindIII fragments or SacI fragments of various sizes were subcloned into the pAU-PS vector. These DNAs were transformed into normal JY745 cells by the above-mentioned method of Okazaki et al. and the acquisition of aureobasidin resistance was examined. As a result, it is revealed that a HindIII-SacI 2.4 kb DNA fragment contains the spaur1^R gene. The restriction enzyme map of this DNA segment containing the aureobasidin resistant gene spaur1^R is shown in Fig. 1. This fragment was cloned into a pUC118 vector (named pUARS2R) and then the DNA

nucleotide sequence was identified (SEQ ID No. 1 in Sequence Listing). From this nucleotide sequence, it is revealed that the spaurl^R gene code for a protein having an amino acid sequence represented by SEQ ID No. 2 in Sequence Listing.

1-e) Cloning of aureobasidin sensitive gene spaurl^S

By the same method as the one employed in the above c), genomic DNA was extracted and purified from normal cells. After partially digesting with HindIII, a genomic library of the normal cells was constructed. An E. coli stock containing this library DNA was spreaded on an LB agar medium containing ampicillin and tetracycline and incubated overnight at 37°C. The colonies thus formed were transferred onto a nylon membrane (Hybond™-N, manufactured by Amersham) and the colony hybridization was performed.

As a probe, the above-mentioned DNA fragment (2.4 kb) obtained by cleaving the spaurl^R gene with HindIII-SacI and labeled with [α -³²p] dCTP by using a random primer DNA labeling kit (manufactured by Takara Shuzo Co., Ltd.) was used. As the results of screening of 5×10^4 colonies, five clones being hybridizable with the probe were obtained. Plasmids were purified from E. coli cells of these five clones. As the result of the cleavage with restriction enzymes, it was found out that all of these clones contained the same DNA fragment of 4.5 kb (named pARN1). The restriction enzyme map of the DNA of 4.5 kb in pARN1 was identical with that of pAR25 shown in

Fig. 10. Therefore, a HindIII-SacI 2.4 kb DNA fragment which was a region containing the *spaur1^s* gene was prepared from pARN1. Then it was cloned into the pAU-PS vector and this plasmid was named pSPAR1.

By using this plasmid pSPAR1, a strain E. coli JM109 was transformed and the transformant thus obtained was named and designated as Escherichia coli JM109/pSPAR1. It has been deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology in accordance with the Budapest Treaty under the accession number FERM BP-4485. This DNA fragment containing the aureobasidin sensitive gene *spaur1^s* had the restriction enzyme map shown in Fig. 1 and the DNA nucleotide sequence thereof was the one represented by SEQ ID NO. 3 in Sequence Listing. Based on this nucleotide sequence, it has been revealed that the *spaur1^s* gene codes for a protein having the amino acid sequence represented by SEQ ID NO. 4 in Sequence Listing and, when compared with the resistant gene *spaur1^r*, the amino acid at the residue 240 has been changed from glycine into cysteine.

Example 2: Cloning of aureobasidin sensitive genes *scaur1* and *scaur2* originating in budding yeast S. cerevisiae

2-a) Separation of aureobasidin resistant mutant of S. cerevisiae

A strain S. cerevisiae DKD5D (mating type a,

genotype leu2-3 112, trp1, his3) having a sensitivity to aureobasidin at a concentration of 0.31 µg/ml was mutagenized with EMS in the same manner as the one employed in the case of Schizo. pombe. Then resistant mutants were separated on an agar plate of a complete nutritional medium YPD (1% of yeast extract, 2% of polypeptone, 2% of glucose) containing 5 µg/ml or 1.5 µg/ml of aureobasidin A. After repeating the mutagenesis several times, 34 mutant clones were obtained. These mutants were resistant to more than 25 µg/ml of aureobasidin A and estimated as having not a multiple drug resistance mutation but a aureobasidin-specific resistance mutation.

2-b) Genetic analysis

Similar to the above-mentioned case of Schizo. pombe, the genetic analysis using the tetrad analysis and the complementation test was performed. As a result, the genes could be classified into two types. These genes regulating aureobasidin sensitivity were named scaur1 and scaur2, the resistant genes isolated from the resistant mutant were named scaur^R1 and scaur^R2, and the sensitive genes isolated from the sensitive wild-type strain were named scaur^s1 and scaur^s2, respectively.

The R94A strain had a gene with dominant mutation (scaur^R1). It has been further clarified that the scaur1 gene is located in the neighborhood of the met14 gene of the eleventh chromosome.

2-c) Preparation of genomic library of aureobasidin resistant strain having aureobasidin resistant gene *scaur1^R*

Genomic DNA was extracted and purified from the aureobasidin resistant strain R94A by the above-mentioned method of P. Philippson et al. The purified genomic DNA (8 µg) was partially digested by treating with 5 U of a restriction enzyme HindIII at 37°C for 10 minutes, deproteinized with phenol/chloroform and precipitated with ethanol. The partially digested DNA thus obtained was electrophoresed on a 0.8% agarose gel and DNA in the region of 3 to 15 kb was extracted. The DNA thus obtained was ligated with a yeast-*E. coli* shuttle vector pWH5 (2 µg) which had been completely digested with HindIII by using a DNA ligation kit and then transformed into *E. coli* HB101. Thus a genomic library was formed. *E. coli* containing this genomic library was cultured in 50 ml of an LB medium containing ampicillin and tetracycline at 37°C overnight. Then a plasmid was recovered and purified from the *E. coli* cells.

2-d) Expression and cloning of aureobasidin resistant gene *scaur1^R*

The above-mentioned genomic library of the R94A strain was transformed into *S. cerevisiae* SH3328 (mating type α, genotype ura3-52, his4, thr4, leu2-3 • 112) in accordance with the method of R.H. Schiestl et al. [Current Genetics, 16, 339 - 346 (1989)]. The transformed cells were spread on a

minimum medium SD plate [0.67% of yeast nitrogen base without amino acids, 2% of glucose, 2% of agar] containing 25 µg/ml of uracil, 35 µg/ml of histidine and 500 µg/ml of threonine. After incubating at 30°C for 3 to 4 days, the colonies thus formed were replicated onto a YPD agar plate containing 1.5 µg/ml of aureobasidin A. A colony thus formed was inoculated into 5 ml of a liquid YPD medium. After incubating at 30°C for 2 days, a plasmid DNA was recovered from the propagated cells by the above-mentioned method of I. Hagan et al. This plasmid was transformed into a yeast again and it was confirmed that the obtained transformant had acquired aureobasidin resistance. This plasmid, which contained a DNA of 3.5 kb, was named pWTCR3. Neither the DNA fragment of 2.0 kb nor the DNA fragment of 1.5 kb obtained by cleaving with HindIII exhibited any aureobasidin resistant activity alone. Thus it is confirmed that the gene is contained in the DNA fragment of 3.5 kb. Fig. 2 shows the restriction enzyme map of this DNA fragment of 3.5 kb containing the aureobasidin resistant gene *scaur1^R*. The HindIII fragments of 1.5 kb and 2 kb were each cloned into pUC118, followed by the determination of the DNA nucleotide sequence (SEQ ID No. 5 in Sequence Listing). From this nucleotide sequence, it has been revealed that the *scaur1^R* gene codes for a protein having an amino acid sequence represented by SEQ ID No. 6 in Sequence Listing.

2-e) Cloning of aureobasidin sensitive gene *scaur1^s*
corresponding to aureobasidin resistant gene
scaur1^R

By the same method as the one employed in the above Example 2-c), genomic DNA was extracted and purified from the parent strain S. cerevisiae DKD5D. After partially digesting with HindIII, the DNA was ligated with pWH5 and transformed into E. coli HB101. Thus a genomic library of the normal cells was formed. An E. coli stock containing this library DNA was spreaded on an LB agar medium containing ampicillin and tetracycline and incubated overnight at 37°C. The colonies thus formed were transferred onto a nylon membrane (Hybond™-N) and the colony hybridization was carried out. As a probe, the DNA fragment of 3.5 kb obtained in the above Example 2-d) and labeled with [α -³²P] dCTP by using a random primer DNA labeling kit (manufactured by Takara Shuzo Co., Ltd.) was used. As the results of screening of 2×10^4 colonies, seven clones being hybridizable with the probe were obtained. Plasmids were purified from E. coli cells of these clones. As the result of the cleavage with restriction enzymes, one of these clones contained a DNA fragment of 3.5 kb. This DNA fragment had the restriction enzyme map of Fig. 2 and thus judged as containing the *scaur1^s* gene. The plasmid containing this DNA fragment was named pSCAR1, while E. coli HB101 having this plasmid introduced therein was named and designated as Escherichia coli HB101/pSCAR1. This

strain has been deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology in accordance with the Budapest Treaty under the accession number FERM BP-4483. The DNA fragment of 3.5 kb obtained by partially digesting pSCARI with HindIII was subcloned into pUC118 and the nucleotide sequence thereof was determined (SEQ ID No. 7 in Sequence Listing). A comparison with the resistant gene indicates that the base at the position 852 has been changed from T into A and, due to this replacement, the amino acid has been converted from phenylalanine into tyrosine (SEQ ID No. 8 in Sequence Listing).

2-f) Preparation of genomic library of aureobasidin resistant strain having aureobasidin resistant gene scaur2^R

A genomic library was prepared from an aureobasidin resistant strain L22-8B by the same method as the one described in Example 2-c). E. coli containing this genomic library was cultured in an LB medium (50 ml) containing ampicillin and tetracycline at 37°C overnight. Then plasmids were recovered and purified from the E. coli cells.

2-g) Expression and cloning of aureobasidin resistant gene scaur2^R

The above-mentioned plasmids originating in the genomic library of the L22-8B strain were transformed into S. cerevisiae SH3328 by the above-mentioned method of R.H. Schiestl. From the transformed

strains, an aureobasidin resistant strain was isolated. Then a plasmid DNA containing the *scaur²*^R gene was recovered from this transformant by the above-mentioned method of I. Hagan et al. This plasmid was transformed into a yeast again and it was confirmed that the transformant had acquired aureobasidin resistance. This plasmid, which contained a DNA of 8.5 kb, was named pSCAR2. Fig. 3 shows the restriction enzyme map of the DNA fragment of 8.5 kb containing this aureobasidin resistant gene *scaur²*^R. *E. coli* HB101 having this plasmid pSCAR2 introduced therein was named and designated as *Escherichia coli* HB101/pSCAR2. This strain has been deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology in accordance with the Budapest Treaty under the accession number FERM BP-4484. By using BamHI, ECORI, HindIII and PstI, DNA fragments of various sizes were prepared and cloned into the pWH5 vector. These plasmids were transformed into *S. cerevisiae* DKD5D in accordance with the above-mentioned method of R.H. Schiestl et al. Then it was examined whether these transformants had acquired aureobasidin resistance or not. As a result, none of the transformants of the DNA fragments was a resistant one. Thus it has been clarified that the DNA fragment of the full length is necessary for the expression of the aureobasidin resistance.

2-h) Isolation of aureobasidin sensitive gene
scaur^s corresponding to aureobasidin
resistant gene scaur^R

An E. coli stock containing the genomic library of Example 2-e) prepared from normal cells of S. cerevisiae DKD5D was spreaded on an LB agar medium containing ampicillin and tetracycline and incubated at 37°C overnight. The colonies thus formed were transferred onto a nylon membrane (Hybond™-N) and the colony hybridization was performed. As a probe the DNA fragment of 8.5 kb obtained in the above Example 2-g) and labeled with [α -³²P] dCTP by using a random primer DNA labeling kit was used. As the results of screening of 2×10^4 colonies, several clones being hybridizable with the probe were obtained. Some of these clones contained a DNA fragment of 4.6 kb while others contained a DNA fragment of 3.9 kb. From the restriction enzyme maps of these DNA fragments, it was found out that these DNA fragments were each a part of the scaur^s gene shown in Fig. 3. These DNA fragments were ligated together to thereby give a scaur^s fragments shown in Fig. 3. The DNA fragment of 8.5 kb thus obtained was subcloned into pUC118 and then the DNA nucleotide sequence was determined (SEQ ID No. 9 in Sequence Listing). Based on the nucleotide sequence of SEQ ID No. 9 in Sequence Listing, the amino acid sequence represented by SEQ ID No. 10 in Sequence Listing was estimated.

Example 3: Gene disruption test on spaurl^s and scauri^s genes

3-a) Gene disruption test on spaurl^s gene

In order to examine whether the aureobasidin sensitive gene spaurl^s is necessary in the cell growth by the gene disruption test, the plasmid pUARS2R prepared in Example 1-d) was first cleaved with BalI and EcoT22I. After eliminating a DNA fragment of 240 bp, the residual DNA fragment was blunted by using a DNA blunting kit (manufactured by Takara Shuzo Co., Ltd.). Then this DNA was ligated with a DNA containing ura4^r gene of 1.7 kb, which had been obtained by excising from a pUC8ura4 plasmid [Mol. Gen. Genet., 215, 81 - 86 (1988)] by cleaving with HindIII and blunting, to thereby give a plasmid pUARS2RBT22::ura4-1 and another plasmid pUARS2RBT22::ura4-6 in which the ura4 DNA had been inserted in the opposite direction.

Both of these disrupted genes were excised from the vector pUC118 by cleaving with SacI and HindIII and ARS2RBT22::ura4-1 and ARS2RBT22::ura4-6 (Fig. 4), which were spaurl^s DNA fragments containing ura4^r, were purified. The purified DNA fragments were transformed into diploid cells Schizo. pombe C525 (h^{90}/h^{90} , ura4-D18/ura4-D18, leu1/leu1, ade6-M210/ade6-M216) by the above-mentioned method of Okazaki et al. and then a transformant was screened on an SD agar plate containing leucine. In the transformant thus obtained, one of a pair of spaurl^s genes on the chromosome had been replaced by the disrupted gene ARS2RBT22::ura4-1.

or ARS2RBT22::ura4-6 introduced thereinto. These cells were allowed to undergo sporulation on a sporulation medium MEA and subjected to the tetrad analysis. As a result, it was found out that two of the four ascospores formed colonies but the residual two spores formed no colony. That is to say, the spores suffering from the replacement of the normal *scaur1^s* gene by the disrupted gene ARS2RBT22::ura4-1 were not propagated. It has been thus revealed that the *scaur1^s* gene is essentially required for the growth of the cells.

3-b) Gene disruption test on *scaur1^s* gene

The plasmid pSCAR1 prepared in Example 2-e) was partially digested with HindIII to thereby give a DNA fragment of 3.5 kb shown in Fig. 2. This DNA fragment was cloned into the HindIII site of pUC119 and the obtained product was named pSCAR3. The obtained pSCAR3 was cleaved with StuI and EcoT22I. After eliminating a DNA fragment of 0.3 kb, the obtained DNA was ligated with a DNA fragment (1.1 kb) of URA3 gene which had been obtained by cleaving a plasmid pYEUrA3 (manufactured by Clontech Laboratories, Inc.) with HindIII and EcoRI and blunting. Thus a plasmid pUSCAR3.ST22::URA3^t and another plasmid pUSCAR3.ST22::URA3A, in which the URA3 gene had been inserted in the opposite direction, were obtained. These disrupted gene were excised in the EcoRI site in the *scaur1^s* gene and the EcoRI site in the pUC119 vector by cleaving with EcoRI. The *scaur1^s* DNA

fragments containing URA3, SCAR3.ST22::URA3⁺ and SCAR3.ST22::URA3A (Fig. 5), were purified. The purified DNA fragments were transformed into diploid cells of S. cerevisiae AOD1 (mating type a/a, genotype ura3-52/ura3-52, leu2-3 112/leu2-3 112, trp1/TRP1, thr4/THR4, his4/HIS4) by the above-mentioned method of R.H. Schiestl and transformants were screened on an SD agar plate containing leucine. The transformants thus obtained were allowed to undergo sporulation on a sporulation medium SP (1% of potassium acetate, 2% of agar) and subjected to the tetrad analysis. As a result, it was found out that two of the four ascospores underwent germination and formed colonies but the residual two spores did not undergo colony formation. That is to say, the spores suffering from the replacement of the *scaur1*^s gene by the disrupted gene were not propagated. It has been thus revealed that the *scaur1*^s gene is essentially required for the growth of the cells.

Example 4. Examination on the expression of aureobasidin sensitive gene *scaur1* by northern hybridization

From a normal strain or a resistant strain of Schizo. pombe, the whole RNAs were extracted and purified by the method of R. Jensen et al. [PROC. NATL. ACAD. SCI. USA, 80, 3035 - 3039 (1983)]. Further, poly(A) RNA was purified by using OligotexTM-dT30 (manufactured by Takara Shuzo Co., Ltd.). The

purified poly(A)⁺RNA (2.5 µg) was separated by the electrophoresis on a 1.2% agarose gel containing formaldehyde and transferred onto a nylon membrane (HybondTM-N). After immobilizing, the hybridization was performed with the use of a HindIII-SacI fragment (2 kb) of the *scaur1^s* gene labeled with [α -³²P]dCTP as a probe. As a result, both of the normal cells and the resistant cells showed a band of the same amount of about 2 kb. In both cases, this amount underwent no change in the logarithmic growth phase and the stationary phase (Fig. 10). Fig. 10 is an autoradiogram showing the results of the northern hybridization wherein mRNAs obtained from cells of a sensitive strain of Schiz. pombe in the logarithmic growth phase (lane 1), cells of a resistant strain in the logarithmic growth phase (lane 2), cells of the sensitive strain in the stationary phase (lane 3) and cells of the resistant strain in the stationary phase (lane 4) are electrophoresed on a 1.2% agarose gel containing formaldehyde.

Example 5: Determination of the activity of *scaur1^s* gene

5-a) Construction of plasmid YEpSCARW3 (Fig. 9) and YEpSCARW1

The plasmid pSCAR1 prepared in Example 2-e) was cleaved with HindIII and a fragment of 2 kb containing the whole ORF was excised. This fragment was inserted

into the HindIII site of a expression-plasmid YEp52 having a promoter Gal10, the expression of which was induced by galactose in a medium. The plasmid having the *scaur1^s* gene which had been inserted in such a direction as to be normally transcribed by the promoter Gal10 was named YEpSCARW3. Fig. 9 shows the structure of this plasmid. Further, the plasmid having the *scaur1^s* gene inserted in the opposite direction was named YEpSCARW1.

5-b) Transformation by plasmids YEpSCARW3 and YEpSCARW1

By using 5 µg portions of the plasmids YEpSCARW3 and YEpSCARW1, the diploid S. cerevisiae cells with the disrupted *scaur1^s* gene prepared in Example 3-b) were transformed. Then transformants were screened on an SD agar plate. These transformants were allowed to undergo sporulation on an SP medium and then subjected to the tetrad analysis. When the expression of the *scaur1^s* gene was induced by using a YPGal medium (1% of yeast extract, 2% of polypeptone, 2% of galactose), the ascospores formed from the diploid cells transformed by YEpSCARW3 all underwent germination while two of the four ascospores formed from the diploid cells transformed by YEpSCARW1 underwent germination but not the remaining two. It is thus conceivable that the cells with the disrupted *scaur1^s* gene have reverted to the normal state by introducing YEpSCARW3 containing the *scaur1^s* gene into these cells. Accordingly, the use of these cells with the

disrupted *scaur1*^s gene as a host makes it possible to determine the activity of normal *aur1*-analogous genes carried by other organisms.

Example 6: Confirmation and cloning of *aur1* and *aur2* genes (*caaur1*, *caaur2*) carried by C. albicans

6-a) Detection of *aur1* gene by the PCR method

Poly(A)⁺RNA was extracted and purified from an aureobasidin sensitive strain C. albicans TIMM0136 by the same method as the one employed in Example 4. By using the poly(A)⁺RNA (5 µg) as a template, a double-stranded cDNA was synthesized on a cDNA synthesizing system Plus (manufactured by Amersham) with the use of an oligo(dT) primer. Mixed primers for PCR corresponding to amino acid sequence regions being common to the amino acid sequences of S. cerevisiae and Schizo. pombe were synthesized on a DNA synthesizer and purified. That is to say, a primer of SEQ ID No. 11 in Sequence Listing corresponding to the region of amino acids at the 184- to 192-positions of SEQ ID No. 4 in Sequence Listing of Schizo. pombe (from the 184- to 192-positions of SEQ ID No. 8 in Sequence Listing of S. cerevisiae) and another primer of SEQ ID No. 12 in Sequence Listing corresponding to the region of amino acids from the 289- to 298-positions of Schizo. pombe (from the 289- to 298-positions of SEQ ID No. 8 in Sequence Listing of S. cerevisiae) were employed.

PCR was performed by using these primers and the above-mentioned cDNA as a template by repeating a cycle comprising treatment at 94°C for 30 seconds, one at 48°C for 1 minute and one at 72°C for 2 minutes 25 times. As a result, a DNA (about 350 bp) being almost the same as S. cerevisiae and Schizo. pombe in length was amplified (Fig. 6). Fig. 6 shows a pattern obtained by carrying out PCR with the use of cDNA of C. albicans (lane 1), cDNA of S. cerevisiae (lane 2) and cDNA of Schizo. pombe (lane 3) as a template, electrophoresing each PCR product on an agarose gel and staining with ethidium bromide.

6-b) Cloning of aurl gene (caaurl) of C. albicans

(i) Genomic DNA was extracted and purified from a strain C. albicans TIMM0136 by the same method as the one described in Example 1-c). After partially digesting with HindIII, the DNA fragment was ligated with a Traplex119 vector which had been completely digested with HindIII and transformed into E. coli HB101. Thus a genomic library of C. albicans was prepared. From this library, a DNA fragment of 4.5 kb containing the aurl gene of C. albicans was cloned by using the DNA fragment of C. albicans obtained by the PCR described in Example 6-a), which had been labeled with [α -³²P]dCTP by using a random primer DNA labeling kit (manufactured by Takara Shuzo Co., Ltd.), as a probe. This DNA fragment had a restriction enzyme map shown in Fig. 7 and the DNA nucleotide sequence thereof is represented by SEQ ID No. 13 in Sequence

Listing. Based on this nucleotide sequence, it was estimated that the caaurl gene coded for a protein having the amino acid sequence represented by SEQ ID No. 14 in Sequence Listing. When compared with the scaur1^s protein, a homology of as high as 53% was observed. A Traplex119 vector having this caaurl gene integrated therein was named pCAAR1, while E. coli HB101 transformed by this plasmid was named and designated as Escherichia coli HB101/pCAAR1. This strain has been deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology in accordance with the Budapest Treaty under the accession number FERM BP-4482.

Next, pCAAR1 was treated with HindIII to thereby give caaurl of 4.5 kb. Further, it was integrated into pTV118 which had been completely digested with HindIII to thereby prepare a plasmid for expressing caaurl. This plasmid was named pTCAAR1.

(ii) Genomic DNA was extracted and purified from a strain C. albicans TIMM1768 [The journal of Antibiotics, 46, 1414-1420(1993)] by the same method as the one described in Example 1-c). After partially digesting with Hind III, the DNA fragment was ligated with a pUC118 vector which had been completely digested with Hind III and transformed into E. coli HB101. Thus a genomic library of C. albicans TIMM1768 was prepared. From this library, a DNA fragment of 4.5 kb containing the aurl gene of C. albicans TIMM1768 was cloned by the colony hybridization with the same

probe as that described in Example 6-b)-(i). This DNA fragment had the same restriction enzyme map as that shown in Fig. 7. Next, a part of the DNA sequence containing a ORF in this DNA fragment was determined. The DNA nucleotide sequence thereof is represented by SEQ ID No. 21 in Sequence Listing. Based on this nucleotide sequence, it was estimated that this gene coded for a protein having the amino acid sequence represented by SEQ ID No. 22 in Sequence Listing. When the amino acid sequence of the caaur1 protein C. albicans TIMM1768 was compared with that of the caaur1 protein of C. albicans TIMM0136, the amino acid sequences of the 1- to 381-positions and the 383- to 423-positions and the 425- to 471-positions of caaur1 protein (SEQ ID No. 14 in Sequence Listing) in C. albicans TIMM0136 were identical with the amino acid sequences of the 2- to 382-positions and the 384- to 424-positions and the 426- to 472-positions, respectively, of caaur1 protein (SEQ ID No. 22 in Sequence Listing) in C. albicans TIMM1768.

However, serines at the 382- and 424-positions of SEQ ID No. 14 in Sequence Listing were replaced with prolines at the 383- and 425-positions of SEQ ID No. 22 in Sequence Listing.

6-c) Cloning of aur2 gene (caaur2) of C. albicans

Genomic DNA of a strain C. albicans TIMM0136 was digested with BamHI and ligated with a pTV118 vector which had been completely digested with BamHI. Then it

was transformed into E. coli HB101 to thereby prepare a genomic library of C. albicans. On the other hand, the DNA fragment containing the scaur^{2s} gene obtained in Example 2-h) was cleaved with HindIII and PstI to thereby give a DNA fragment of 1.2 kb. This DNA fragment was labeled with [α -³²P]dCTP by using a random primer DNA labeling kit. By using this labeled DNA fragment as a probe, the above-mentioned C. albicans genomic library was screened by the colony hybridization. Thus a plasmid containing a DNA fragment of 8.3 kb was obtained. A part of the DNA sequence upstream of the BamHI site of this DNA fragment was determined (SEQ ID No. 15 in Sequence Listing). Based on this sequence, an amino acid sequence represented by SEQ ID No. 16 in Sequence Listing was estimated. It corresponded to the amino acid sequence of the 1230- to 1309-positions of the amino acid sequence of the scaur² gene (SEQ ID No. 10), having a homology of as high as 77%. Since this DNA fragment lacked a part of the C-end, the genomic library prepared in Example 6-b) was further screened by using this DNA fragment as a probe. Thus a DNA fragment of 6.5 kb having the C-terminal part was obtained. Fig. 8 shows the restriction enzyme map of the DNA region containing the caaur² gene thus clarified.

A pTV118 vector having the above-mentioned caaur² gene of 8.3 kb integrated therein was named pCAAR2N, while E. coli HB101 transformed by this plasmid was

named and designated as Escherichia coli HB101/pCAAR2N. This strain has been deposited at National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology in accordance with the Budapest Treaty under the accession number FERM BP-4481.

Example 7: Preparation of antibody against protein coded for by scaur1^s gene and staining of S. cerevisiae cells and detection of said protein by using this antibody.

7-a) Preparation of antibody

SCAR1-1 (SEQ ID No. 19 in Sequence Listing) comprising a peptide corresponding to the amino acids at the residue 103 to 113 in the amino acid sequence of SEQ ID No. 8 in Sequence Listing having cysteine added to the N-end thereof and SCAR1-2 (SEQ ID No. 20 in Sequence Listing) comprising a peptide corresponding to the amino acids at the residue 331 to 348 in the amino acid sequence of SEQ ID No. 8 having cysteine added to the N-end thereof were synthesized by the Fmoc solid phase synthesis method and purified by reverse phase HPLC. Thus 10 mg portions of these peptides were obtained. To the N-terminal cysteine of each of these synthetic peptides, KLH was bound as a carrier protein. By using this binding product as an antigen, a rabbit was immunized and an antiserum was obtained. This antiserum was further purified on an affinity column prepared by binding the synthetic

peptide employed as the antigen to an agarose gel.

This a polyclonal antibody being specific for the synthetic peptide was prepared.

7-b) Staining of S. cerevisiae cells with antibody

A strain S. cerevisiae ATCC 9763 was cultured in a YNBG medium [0.67% of yeast nitrogen base (manufactured by Difco), 2% of glucose] to thereby give a suspension of a concentration of 3×10^7 cells/ml. To 1 ml of this cell suspension were added 0.11 ml of a 1 M phosphate buffer (pH 6.5) and 0.17 ml of 37% formaldehyde. After slowly stirring at room temperature for 1 hour, the cells were harvested by centrifugation and then suspended in 20 ml of an SS buffer (1 M of sorbitol, 0.2 % of β -mercaptoethanol, 0.1 M phosphate buffer, pH 7.5) containing 20 μ g/ml of Zymolyase 20T. After treating at 30°C for 1 hour, the cells were harvested, washed with the SS buffer, suspended in 1 ml of the SS buffer containing 0.1% of Triton X-100 and then allowed to stand for 10 minutes. This cell suspension was placed on a slide glass which had been coated with poly(L-lysine) and allowed to stand for 10 minutes. Next, a PBS solution containing 1% of albumin (BSA) was dropped thereinto. After allowing to stand at room temperature for 15 minutes, the excessive liquid was removed and then a PBS solution containing BSA containing 0.02 mg/ml of the antiSCAR1-1 antibody was dropped thereinto. After allowing to stand at room temperature for 60 minutes and washing with PBS containing BSA three times,

antirabbit IgG antibody labeled with FITC (antibody concentration 0.02 mg/ml) was layered over and allowed to stand at room temperature for 1 hour. After washing with a PBS solution containing BSA, a small amount of a mountain solution, which was a solution prepared by dissolving 0.1 g of p-phenylenediamine in 10 ml of CBS (150 mM of NaCl, 50 mM of CHES, pH 9.5), adjusting the pH value to 9.0 with 10 N NaOH and further adding 90 ml of glycerol, was layered over. Then a cover glass was placed thereon to thereby give a specimen. This specimen was observed under a fluorescence microscope to thereby examine the intracellular distribution of the scaur1 protein. As a result, it was found out that this protein was distributed all over the cells.

7-c) Detection of protein coded for by scaur1 gene by using antibody

The plasmid YEpSCARW3 prepared in Example 5-a) was introduced into a normal haploid S. cerevisiae SH3328 to thereby give a transformant. This transformant was cultured in a YPGal medium or a YPD medium and the cells were harvested by centrifugation. The cells thus obtained were suspended in a buffer (1% of Triton X-100, 1% of SDS, 20 mM of Tris-HCl, pH 7.9, 10 mM of EDTA, 1 mM of DTT, 1 mM of PMSF). Further, glass beads were added thereto to disrupt the cells by vigorous vortex. Then an SDS loading solution was added thereto and the protein was denatured by treating at 95°C for 5 minutes. After centrifuging, a part of the obtained supernatant was subjected to SDS-PAGE and the protein

thus separated was transferred onto an Immobilon membrane (manufactured by MILLIPORE). This Immobilon membrane was treated with Block Ace (manufactured by Dainippon Pharmaceutical Co., Ltd.). Then the antiSCAR1-2 antibody prepared in 7-a) was reacted therewith as a primary antibody. After washing, antirabbit IgG antibody labeled with peroxidase was reacted therewith as a secondary antibody and the mixture was thoroughly washed. Next, it was color-developed with diaminobenzidine and a band of the scaur1 protein was detected. Fig. 11 shows the results.

Fig. 11 shows the results of the detection of the protein prepared from the cells incubated in the YPD medium (lane 1) and the protein prepared from the cells incubated in the YPGal medium (lane 2), each subjected to SDS-PAGE, by using the antiSCAR1-2 antibody. The cells incubated in the YPGal medium, of which scaur1 gene had been induced, showed a specific band.

[Effects of the Invention]

According to the present invention, a novel protein regulating aureobasidin sensitivity and a gene coding for the protein, i.e., a gene regulating aureobasidin sensitivity are provided. These substances are useful in the diagnosis and treatment for diseases caused by organisms having the above-mentioned gene, such as mycoses. The present invention further provides an

antisense DNA and an antisense RNA of this gene, a nucleic acid probe being hybridizable with the gene, a process for detecting the gene by using this nucleic acid probe, a process for producing a protein regulating aureobasidin sensitivity by using a transformant having the gene introduced thereinto, an antibody for the protein and a process for detecting the protein by using this antibody. They are also useful in the diagnosis and treatment of diseases including mycoses.

Sequence Listing

SEQ ID NO : 1

SEQUENCE LENGTH : 2385

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION :

AAGCTTTT GCCTCTGAA AAGTCGTT CTCGAATTGG TTTTGAGG AAAAGCAACT 60
TAATAAACTA ATTATATTAT ATATAATTAG CAATTTATA AAAAAAATAA AAAAATACCC 120
CTGATTGCTG GCAACTGTGA GCTGAACATT GGTTAATCGG TCCATCTTT TTTAAATATT 180
TTACATCGCT ACITTTAAGT CCTTGACACT TGCATTTAAT AGCTACTTTC TTTCCTTCAT 240
AAAAATTCCCT TTTTTTCCT TTAGTTTCC CGTTAATTCC TTACGAAATT TTTTCTGTAC 300
GCTCCCTTT TTIACTGTGA TAATTCCTTG AAGCAATGTC TGCTCTTCG ACCTTAAAAA 360
AGCGCCTTGC TCCGTGTAAAC CGACCATCCC AAATACAAGTT CGAAACAAACC TTAAACCCCTA 420
TCCCTACATT TCGTTTCTA CGCAATACGA AATGGTCATG GACACATTG CAATATCTGT 480
TTCTAGCAGG TAATTGATT TTGCTTGTG TTGTCATTGA ATCTCCTGGA TTCTGGGGGA 540
AATTGGCAT TCCCTGTCTT TTGCCCATTG CGTTGACCGT TCCTTTAACAA CCCCAAAATT 600
TTTTCTGCC CATTGTTATC ATCACCTGGG CAATTTATT TTACTCTTGT AGCTTTATTC 660
CAGAACGGTC CGCTCCACCC ATATGGGTTTC GTCTTTACCC CACACTTGAA AATATTCTT 720
ATGGCTCTAA TCTTTCTAGT CTTCTCTGAA AAACCACGCA TAGCATCCTT GATATTCTG 780
CCTGGGTTCC ATATGGAGTC ATGCATTATT CGGCTCCTT TATCATTCA TTTATTCTT 840
TCATCTTGC ACCTCCTGGA ACTCTTCCAG TTTGGGCTCG AACTTTGGT TATATGAATT 900
TATTTGGTGT TCTTATCCAA ATGGCTTCC CGTGTCTCC TCCTTGGTAT GAAAATATGT 960
ATGGTTTAGA ACCTGCCACG TATGCACTAC GTGGCTCTCC TGGTGGATTG GCGGGTATTG 1020
ATCCCTCTCTT CGGCACTAGC ATTTACACTG ATTGTTTTC TAACTCTCCG GTTGTTTTG 1080
GTGCCCTTCC ATCTCTTCAC GCTGGATGGG CCATGCTGGA AGCACTTTTC CTTCGGCATG 1140
TCTTTCTCG ATACCGCTTC TGCTTTATG GATATGTTCT ATGGCTTGC TGGTGTACTA 1200
TGTACCTTAC CCACCACTAC TTTGTAGATT TGGTCGGCGG TATGTGTTA GCTATTATAT 1260

GCTTCGTTT TGCTCAAAAG CTACGCCCTCC CACAGTTGCA AACTGGTAAA ATCCTTCGTT 1320
 GGGAAATACGA GTTTGTTATC CACGGTCATG GTCTTCCGA AAAAACCGAGC AACTCCTTGG 1380
 CTCGTACCGG CAGCCCCATAC TTACTTGAA GGGATTCTTT TACTCAAAAC CCTAATGCAG 1440
 TAGCCCTCAT GAGTGGTCTT AACAAATATGG AACTTGCTAA CACCGATCAT GAATGGTCCG 1500
 TGGGTTCATC ATCACCTGAG CCGTTACCTA GTCCTGCTGC TGATTGATT GATCGTCCTG 1560
 CCAGTACAC TTCCCTCCATC TTTGATGCAA GTCATCTTCC TTAAATCAAC GTGCTTAAG 1620
 AATATATTTC CAAAAGCTAC ATGATACATT GACTAGAATC GGTTTGATT ATAGTGGTAT 1680
 TGGAATGATG TTGTTCATTG TGTTTTAA CTGTTAATCT GACATCCATT GAGTCATTCT 1740
 TTACAATTG TAAAATTAAT TTGTATCACT AATTTGAAG GAAGCTATTT TGGTATTAAT 1800
 ACCGCTTTG GTCTCCACTT CCTTTTCGAA ACTCTTAACA GCGATTAGGC CGGGTATCTT 1860
 CCAGTGTGAT GTATAGGTAT TTGTCGTTT TTTATCATT CCGTTAATAA AGAACTCTT 1920
 TATCCAGCTT CTTACACTGT CAACTGTTGT GAAAGGAACA CATTAGAAT TTCATTTCC 1980
 TTATTTGTTG TGATTTAAAT CGTTGACAT AATTTAAAT TTGGTTGAA ATGTGTGTGA 2040
 GAAGGCTTGT TTTATTCAATT TAGTTTATTG CTGGTTGCA CGAAAATCCA GAACGGAGCA 2100
 TTAAATGTAAT CCTTTTTTAT TCTGTAAAGC GTTTTATAC AAATGTTGGT TATACGTTTC 2160
 TAAAATAAGA ATATTGTTAT AATAATATAG TTTTTCTAT CATTGTTAC ACACACTAAA 2220
 GAGACATTAA GGATAAGCAA ATGTGTTAAA ATGATAATAT ATTTGGAAA CATTATAAA 2280
 GAAATTAAGC ACCTTGACT AACTACATT TTGTTTTT CCTAAGGAAA ACTGTATAGT 2340
 TATACACGCG AGCTGTATTC ACTTCCATTG TAGTGACTTG AGCTC 2385

SEQ ID NO : 2

SEQUENCE LENGTH : 422

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Ser Ala Leu Ser Thr Leu Lys Lys Arg Leu Ala Ala Cys Asn

1

5

10

15

Arg Ala Ser Gln Tyr Lys Leu Glu Thr Ser Leu Asn Pro Met Pro
 20 25 30
 Thr Phe Arg Leu Leu Arg Asn Thr Lys Trp Ser Trp Thr His Leu
 35 40 45
 Gln Tyr Val Phe Leu Ala Gly Asn Leu Ile Phe Ala Cys Ile Val
 50 55 60
 Ile Glu Ser Pro Gly Phe Trp Gly Lys Phe Gly Ile Ala Cys Leu
 65 70 75
 Leu Ala Ile Ala Leu Thr Val Pro Leu Thr Arg Gln Ile Phe Phe
 80 85 90
 Pro Ala Ile Val Ile Ile Thr Trp Ala Ile Leu Phe Tyr Ser Cys
 95 100 105
 Arg Phe Ile Pro Glu Arg Trp Arg Pro Pro Ile Trp Val Arg Val
 110 115 120
 Leu Pro Thr Leu Glu Asn Ile Leu Tyr Gly Ser Asn Leu Ser Ser
 125 130 135
 Leu Leu Ser Lys Thr Thr His Ser Ile Leu Asp Ile Leu Ala Trp
 140 145 150
 Val Pro Tyr Gly Val Met His Tyr Ser Ala Pro Phe Ile Ile Ser
 155 160 165
 Phe Ile Leu Phe Ile Phe Ala Pro Pro Gly Thr Leu Pro Val Trp
 170 175 180
 Ala Arg Thr Phe Gly Tyr Met Asn Leu Phe Gly Val Leu Ile Gln
 185 190 195
 Met Ala Phe Pro Cys Ser Pro Pro Trp Tyr Glu Asn Met Tyr Gly
 200 205 210
 Leu Glu Pro Ala Thr Tyr Ala Val Arg Gly Ser Pro Gly Gly Leu
 215 220 225
 Ala Arg Ile Asp Ala Leu Phe Gly Thr Ser Ile Tyr Thr Asp Cys

230 235 240
Phe Ser Asn Ser Pro Val Val Phe Gly Ala Phe Pro Ser Leu His
245 250 255
Ala Gly Trp Ala Met Leu Glu Ala Leu Phe Leu Ser His Val Phe
260 265 270
Pro Arg Tyr Arg Phe Cys Phe Tyr Gly Tyr Val Leu Trp Leu Cys
275 280 285
Trp Cys Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Val
290 295 300
Gly Gly Met Cys Leu Ala Ile Ile Cys Phe Val Phe Ala Gln Lys
305 310 315
Leu Arg Leu Pro Gln Leu Gln Thr Gly Lys Ile Leu Arg Trp Glu
320 325 330
Tyr Glu Phe Val Ile His Gly His Gly Leu Ser Glu Lys Thr Ser
335 340 345
Asn Ser Leu Ala Arg Thr Gly Ser Pro Tyr Leu Leu Gly Arg Asp
350 355 360
Ser Phe Thr Gln Asn Pro Asn Ala Val Ala Phe Met Ser Gly Leu
365 370 375
Asn Asn Met Glu Leu Ala Asn Thr Asp His Glu Trp Ser Val Gly
380 385 390
Ser Ser Ser Pro Glu Pro Leu Pro Ser Pro Ala Ala Asp Leu Ile
395 400 405
Asp Arg Pro Ala Ser Thr Thr Ser Ser Ile Phe Asp Ala Ser His
410 415 420
Leu Pro

SEQ ID NO : 3

SEQUENCE LENGTH : 2385

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION :

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CTGATTGCTG GCAACTGTGA GCTGAACATT GGTTAACCGG TCCATCTTT TTTAAATATT 180
TTACATCGCT ACCTTTAAGT GCTTGACACT TGCAATTAAAT AGCTACTTTC TTTCCTTCAT 240
AAAAATTCCT TTTTTTCCT TTAGTTTCC GGTTAATTCC TTACGAAATT TTTTCGTAC 300
GGTTCCCTT TTTACTCTGA TAATTCTTG AAGCAATGTC TGCTCTTCG ACCTTAAAAA 360
AGCCCCTTGC TGGGTGTAAC CGAGCATCCC AATACAAGTT GGAAAGAAGC TTAAACCGTA 420
TGCCTACATT TCGTTTGCAT CGCAATAGGA AATGGTCATG GACACATTG CAATATGCT 480
TTCTAGGAGG TAATTGATT TTTGCTTGTG TTGTGATTGA ATCTCCTGGG TTCTGGGGGA 540
AATTGGCAT TGCCTGTCTT TTGGCCATTG CGTTGACCGT TCCTTTAACG CGCCAAATTT 600
TTTTCCCTGC CATTGTTATC ATCACCTGGG CAATTTATT TTACCTTGT AGGTTTATTG 660
CAGAACGGTG CGGTCCACCC ATATGGGTC GTGTTTACG CACACTTGA AATATTCTT 720
ATGGCTCAA TCTTCTAGT CTTCCTGCA AAACCACGCA TAGCATCCTT GATATTTGG 780
CCTGGGTCTCC ATATGGACTC ATGGATTATT CGGCTCCTTT TATCATTGCA TTTATGTTT 840
TCATGTTGCA ACCTCCTGCA ACTCTTCCAG TTTGGGCTCG AACTTTGGT TATATGAATT 900
TATTGGTGT TCTTATCCAA ATGGCTTCC CGTCTTCTCC TCCTTGCTAT GAAAATATGT 960
ATGGTTTAGA ACCTGCCACG TATGCAGTAC GTGGCTCTCC TGGTGGATTG GCCCGTATTG 1020
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TGTTTCCCTG ATACCCCTTC TGCTTTATG CATATGTTCT ATGGCTTGC TGGTGTACTA 1200
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GCTTCCGTTT TGCTCAAAAG CTACGCCCTCC CACAGTTGCA AACTGCTAAA ATCCTTCGTT 1320
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CTCGTACCGG CAGCCCCATAC TTACTTGGAA GGGATTCTTT TACTCAAAAC CCTAATGGAG 1440

TAGCCTTCAT GAGTGGTCTT AACAAATATGG AACCTGCTAA CACCGATCAT GAATGGTCCG 1500
TGGGTTCATC ATCACCTGAG CCGTTACCTA GTCCCTGCTGC TGATTTGATT GATCGTCCTG 1560
CCAGTACCAAC TTCCCTCCATC TTTGATGCAA GTCATCTTCC TTAAATCAAC GTGCTTAAG 1620
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TTACAATTTG TAAAATTAAT TTGTATCACT AATTTGAAG GAAGCTATTT TGGTATTAAT 1800
ACCGCTTTG GTCTCCACTT CCTTTGAA ACTCTTAACA GCGATTAGGC CGGGTATCTT 1860
CCAGTGTGAT GTATAGGTAT TTGTCGTTT TTTATCATT CCCTTAATAA AGAACTCTTT 1920
TATCCAGCTT CTTACACTGT CAACTGTTGT GAAAGGAACA CATTAGAAT TTCATTTCC 1980
TTATTTGTTG TGATTTAAAT CGTTTGACAT AATTTAAAT TTGGTTGAA ATGTGTGTGA 2040
GAAGCCCTGT TTATTCATT TAGTTTATTG CTTGTTGCA CGAAAATCCA GAACGGAGGA 2100
TTAATGTAAT CCTTTTTTAT TGTGAAAGC GTTTTATAC AAATGTTGGT TATAGGTTTC 2160
TAAAATAAGA ATATTTGTTAT AATAATATAG TTTTTCTAT CATTGTTAC ACACACTAAA 2220
GAGACATTAAGC AAAGGTTAAATGTTGTTAAATGATAATAT ATTTGGAAA GATTATTTAA 2280
GAAATTAAAGC AGCTTTGACT AACCTACATT TTGTTTTTTT CCTAAGGAAA ACTGTATAGT 2340
TATACACGGG AGCTGTATTG ACTTCCATTG TAGTGACTTG AGCTC 2385

SEQ ID NO : 4

SEQUENCE LENGTH : 422

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Ser Ala Leu Ser Thr Leu Lys Lys Arg Leu Ala Ala Cys Asn

1 5 10 15

Arg Ala Ser Gln Tyr Lys Leu Glu Thr Ser Leu Asn Pro Met Pro

20 25 30

Thr Phe Arg Leu Leu Arg Asn Thr Lys Trp Ser Trp Thr His Leu

35	40	45
Gln Tyr Val Phe Leu Ala Gly Asn Leu Ile Phe Ala Cys Ile Val		
50	55	60
Ile Glu Ser Pro Gly Phe Trp Gly Lys Phe Gly Ile Ala Cys Leu		
65	70	75
Leu Ala Ile Ala Leu Thr Val Pro Leu Thr Arg Gln Ile Phe Phe		
80	85	90
Pro Ala Ile Val Ile Ile Thr Trp Ala Ile Leu Phe Tyr Ser Cys		
95	100	105
Arg Phe Ile Pro Glu Arg Trp Arg Pro Pro Ile Trp Val Arg Val		
110	115	120
Leu Pro Thr Leu Glu Asn Ile Leu Tyr Gly Ser Asn Leu Ser Ser		
125	130	135
Leu Leu Ser Lys Thr Thr His Ser Ile Leu Asp Ile Leu Ala Trp		
140	145	150
Val Pro Tyr Gly Val Met His Tyr Ser Ala Pro Phe Ile Ile Ser		
155	160	165
Phe Ile Leu Phe Ile Phe Ala Pro Pro Gly Thr Leu Pro Val Tyr		
170	175	180
Ala Arg Thr Phe Gly Tyr Met Asn Leu Phe Gly Val Leu Ile Gln		
185	190	195
Met Ala Phe Pro Cys Ser Pro Pro Trp Tyr Glu Asn Met Tyr Gly		
200	205	210
Leu Glu Pro Ala Thr Tyr Ala Val Arg Gly Ser Pro Gly Gly Leu		
215	220	225
Ala Arg Ile Asp Ala Leu Phe Gly Thr Ser Ile Tyr Thr Asp Gly		
230	235	240
Phe Ser Asn Ser Pro Val Val Phe Gly Ala Phe Pro Ser Leu His		
245	250	255

Ala Gly Trp Ala Met Leu Glu Ala Leu Phe Leu Ser His Val Phe
260 265 270
Pro Arg Tyr Arg Phe Cys Phe Tyr Gly Tyr Val Leu Trp Leu Cys
275 280 285
Trp Cys Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Val
290 295 300
Gly Gly Met Cys Leu Ala Ile Ile Cys Phe Val Phe Ala Gin Lys
305 310 315
Leu Arg Leu Pro Gin Leu Gin Thr Gly Lys Ile Leu Arg Trp Glu
320 325 330
Tyr Glu Phe Val Ile His Gly His Gly Leu Ser Glu Lys Thr Ser
335 340 345
Asn Ser Leu Ala Arg Thr Gly Ser Pro Tyr Leu Leu Gly Arg Asp
350 355 360
Ser Phe Thr Gin Asn Pro Asn Ala Val Ala Phe Met Ser Gly Leu
365 370 375
Asn Asn Met Glu Leu Ala Asn Thr Asp His Glu Trp Ser Val Gly
380 385 390
Ser Ser Ser Pro Glu Pro Leu Pro Ser Pro Ala Ala Asp Leu Ile
395 400 405
Asp Arg Pro Ala Ser Thr Thr Ser Ser Ile Phe Asp Ala Ser His
410 415 420
Leu Pro

SEQ ID NO : 5

SEQUENCE LENGTH : 2340

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION :

TTCTTTCTG TCAAAGAATA ATAAAGTGC CATCAGTGTT CATAATTGTT ACAAAAGTGGT 60
TTTCTGATTT GGTACTACTG CAGAGGCGTA TTTTTGCTT CAGTTACCAT AGCGTAAGAA 120
CACTAGCGAC TTTTGTTCGT GAACCAACAG ACTAGGATT CTACTGCTAC ATCTCTTAGG 180
TAGTTGCTTA GTCCGATCGC TCACCTTTGG TTGTTGTTAA GTACTTCATA AGTTTATCCT 240
TTTCCTTTT CACACTGAGC TACTTTGGT ATAGCTTTG GCCCAAGGAT CTTGAATT 300
TCTCCAAAAG TACTTTATTT TATATCCTAC AGGTTGCGGT TTTCATATT TAAAAAGCTT 360
TTAATCATT CCTTTGCCGA TGCCAAACCC TTTTTCGAGA TGTTTCTAT CAGAGAGACC 420
TCCAAACTGC CATGTAGCCG ATTTAGAAAC AAGTTTAGAT CCCCATCAA CGTTGTTGAA 480
GGTGCAAAAAA TACAAACCCG CTTAACCGA CTGGGTGCAT TACATCTTCT TCGGATCCAT 540
CATGCTGTTT GTGTTCATTA CTAATCCCAGC ACCTTGGATC TTCAAGATCC TTTTTTATTG 600
TTTCTTGGGC ACTTTATTCA TCATTCCAGC TACGTACAG TTTTCTTCA ATGCCCTGCC 660
CATCCTAACCA TGGGTGGCGC TGTATTCAC TTCATCGTAC TTTCCAGATG ACCGCAGGCC 720
TCCTATTACT GTCAAAGTGT TACCAAGGGT GGAAACAATT TTATACGGCG ACAATTAAAG 780
TGATATTCTT GCAACATCGA CGAATTCTT TTTGGACATT TTAGCATGGT TACCGTACGG 840
ACTATTCAT TATGGGGCCC CATTGTCGT TGCTGCCATC TTATTCGTAT TTGGTCCACC 900
AACTGTTTTG CAAGGTTATG CTTTGCATT TGTTATATG AACCTGTTG GTGTTATCAT 960
GCAAAATGTC TTTCCAGCCG CTCCCCATG GTATAAAATT CTCTATGGAT TGCAATCACC 1020
CAACTATGAT ATGCATGGCT CGCCTGGTGG ATTAGCTAGA ATTGATAAGC TACTCGGTAT 1080
TAATATGTAT ACTACAGCTT TTCAAAATTC CTCCGTCAATT TTGGTGTGCTT TTCCCTTCACT 1140
GCATTCCGGG TGTGCTACTA TCGAAGGCCCT GTTTTCTGT TATTGTTTTC CAAAATTGAA 1200
GCCCTTGTGTT ATTGCTTATG TTGCTGGTT ATGGTGTCA ACTATGTATC TGACACACCA 1260
TTATTTGTA GACCTTATGG CAGGTTCTGT GCTGTACATC GTTATTTCC AGTACACACAA 1320
GTACACACAT TTACCAATTG TAGATACATC TCTTTTGC AGATGGTCAT ACACCTCAAT 1380
TGAGAAATAC GATATATCAA AGAGTGATCC ATTGGCTGCA GATTCAAACG ATATCGAAAG 1440
TGTCCCTTTG TCCAACCTGG AACTTGACTT TGATCTTAAT ATGACTGATG AACCCAGTGT 1500
AAGCCCTTCG TTATTTGATG GATCTACTTC TGTTTCTCGT TCGTCCGCCA CGTCTATAAC 1560
GTCACTAGGT GTAAAGAGGG CTTAACATGAGT ATTTTATCTG CAATTACCGA TACGGTTGGT 1620

CTTATGTAGA TACATATAAA TATATATCTT TTTCTTCCTT TTTCTTAGTC AGGATTGTCG 1680
TTTAGCATAA TATACATGTA CTTTATTTAA TCACATACCA CTGATTATCT TTAGAATTAA 1740
ATAAAATTTT GAAATAAAATG GGTGGCTTTT AATGGGTGCT ATGTTAAGTG AGGCTTTAG 1800
AATGCTCTTC CTGCTTGTT TATTATATGT GTATGAAAGA TATGTATGTA TTTACATGTC 1860
TTTGTAGCGT CCCCAGTCAA AACCTGTGCG CTATACCTAA ATGGATTGAT AATCTTCATT 1920
CACTAATTCT AAAATAGACT TCTTCCCCAA AGAACGGTGT AACGATGAGG CTCTATCCAG 1980
CTGCTTATCT AAATCAACTT TAACGATGGA TGATCTTATG ACACGGGGAT CTTTCTTAA 2040
AGTTCTTAGA ATTTCAGACT GTACCGCAGC TGATGAATCA AACAGCATTA AAAAGTGATA 2100
TGCTCGAAAA TGTTTTCTT GGTCTTCCTT CATTATTTA GGAAGATACC TTATGCCAT 2160
GGGTACAATG TCCCTCACCA CACCTCTGTT TTGAATAATC AGTTTCCGA TTGTGGAAGA 2220
CAATTCTTT GCTTCCAACCT TTGGCGCATT CGAGTTGGTT ATGCGAACAA GTCCGATCAG 2280
CTCATAAAAGC ATCTTACTGAA AAAGGGTGGT TTTGCCTTAT TCTTTCCCTCT GTTGAAGCTT 2340

SEQ ID NO : 6

SEQUENCE LENGTH : 401

SEQUENCE TYPE : amino acid

STRANDED : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Ala Asn Pro Phe Ser Arg Trp Phe Leu Ser Glu Arg Pro Pro

1 5 10 15

Asn Cys His Val Ala Asp Leu Glu Thr Ser Leu Asp Pro His Gln

20 25 30

Thr Leu Leu Lys Val Gln Lys Tyr Lys Pro Ala Leu Ser Asp Trp

35 40 45

Val His Tyr Ile Phe Leu Gly Ser Ile Met Leu Phe Val Phe Ile

50 55 60

Thr Asn Pro Ala Pro Trp Ile Phe Lys Ile Leu Phe Tyr Cys Phe

65	70	75
Leu Gly Thr Leu Phe Ile Ile Pro Ala Thr Ser Gln Phe Phe Phe		
80	85	90
Asn Ala Leu Pro Ile Leu Thr Trp Val Ala Leu Tyr Phe Thr Ser		
95	100	105
Ser Tyr Phe Pro Asp Asp Arg Arg Pro Pro Ile Thr Val Lys Val		
110	115	120
Leu Pro Ala Val Glu Thr Ile Leu Tyr Gly Asp Asn Leu Ser Asp		
125	130	135
Ile Leu Ala Thr Ser Thr Asn Ser Phe Leu Asp Ile Leu Ala Trp		
140	145	150
Leu Pro Tyr Gly Leu Phe His Tyr Gly Ala Pro Phe Val Val Ala		
155	160	165
Ala Ile Leu Phe Val Phe Gly Pro Pro Thr Val Leu Gln Gly Tyr		
170	175	180
Ala Phe Ala Phe Gly Tyr Met Asn Leu Phe Gly Val Ile Met Gln		
185	190	195
Asn Val Phe Pro Ala Ala Pro Pro Trp Tyr Lys Ile Leu Tyr Gly		
200	205	210
Leu Gln Ser Ala Asn Tyr Asp Met His Gly Ser Pro Gly Gly Leu		
215	220	225
Ala Arg Ile Asp Lys Leu Leu Gly Ile Asn Met Tyr Thr Thr Ala		
230	235	240
Phe Ser Asn Ser Ser Val Ile Phe Gly Ala Phe Pro Ser Leu His		
245	250	255
Ser Gly Cys Ala Thr Met Glu Ala Leu Phe Phe Cys Tyr Cys Phe		
260	265	270
Pro Lys Leu Lys Pro Leu Phe Ile Ala Tyr Val Cys Trp Leu Trp		
275	280	285

Trp Ser Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Met
 290 295 300
 Ala Gly Ser Val Leu Ser Tyr Val Ile Phe Gln Tyr Thr Lys Tyr
 305 310 315
 Thr His Leu Pro Ile Val Asp Thr Ser Leu Phe Cys Arg Trp Ser
 320 325 330
 Tyr Thr Ser Ile Glu Lys Tyr Asp Ile Ser Lys Ser Asp Pro Leu
 335 340 345
 Ala Ala Asp Ser Asn Asp Ile Glu Ser Val Pro Leu Ser Asn Leu
 350 355 360
 Glu Leu Asp Phe Asp Leu Asn Met Thr Asp Glu Pro Ser Val Ser
 365 370 375
 Pro Ser Leu Phe Asp Gly Ser Thr Ser Val Ser Arg Ser Ser Ala
 380 385 390
 Thr Ser Ile Thr Ser Leu Gly Val Lys Arg Ala
 395 400

SEQ ID NO : 7

SEQUENCE LENGTH : 2340

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION :

TTTCTTCCTG TCAAAGAATA ATAAAGTGC	CATCA	GTGTT CATATTGTT ACAAA	GTGGT 60
TTTCTGATTG GGTACTACTG CAGAGCGTA	TTTTTGCTT CAGTTACCAT AGCGTAAGAA 120		
CACTAGCGAC TTTGTTCGT GAACCAACAG	AGTAGGATT	CTACTGCTAC ATCTCTTAGG 180	
TAGTTGGTTA GTCCGATCGC TCAC	TTTTGG TTGTTGTTAA GTACTTCATA AGTTTATCCT 240		
TTTCCTTTT CACACTGAGC TACTTGGGT	ATAGCTTTG GCCCAAGGAT CTTGAATT	300	

TCTCCAAAAG TACTTTATT TATATCCTAC AGGTTGCCTT TTTCATATT TAAAAAGCTT 360
TTTAATCATT CCTTTGCCTA TGGCAAACCC TTTTCGAGA TGTTTCTAT CAGAGAGACC 420
TCCAAACTGC CATGTAGCCG ATTTAGAAC AAGTTAGAT CCCCATCAA CGTTGTTGAA 480
GGTGCAAAAA TACAAACCCG CTTTAAGCGA CTGGGTGCAT TACATCTCT TGGGATCCAT 540
CATGCTGTT GTGTTCATTA CTAATCCGC ACCTTGGATC TTCAAGATCC TTTTTATTG 600
TTTCTTGGGC ACTTTATTCA TCATTCAGC TACGTACAG TTTTCCTCA ATGCCTGCC 660
CATCCTAACCA TGGGTGGCGC TGTATTCAC TTCATCGTAC TTTCCAGATG ACCGCAGGCC 720
TCCTATTACT GTCAAAGTGT TACCAGCGT GGAAACAATT TTATACGGCG ACAATTAAAG 780
TGATATTCTT GCAACATCGA CGAACATCCTT TTTGGACATT TTAGCATGGT TACCGTACGG 840
ACTATTCAT TTTGGGGCCC CATTGTCGT TGCTGCCATC TTATCGTAT TTGGTCCACC 900
AACTGTTTG CAAGGTTATG CTTTGCAATT TGGTTATATG AACCTGTTG GTGTTATCAT 960
GCAAAATGTC TTTCCAGCCG CTCCCCATG GTATAAAATT CTCTATGGAT TGCAATCAGC 1020
CAACTATGAT ATGCATGGCT CGCCTGGTGG ATTAGCTAGA ATTGATAAGC TACTCGGTAT 1080
TAATATGAT ACTACAGCTT TTTCAAATT CTCGGTCATT TTCGGTGCTT TTCTTCACT 1140
GCATTCCGGG TGTGCTACTA TGGAAAGCCCT GTTTTCTGT TATTGTTTC CAAAATTGAA 1200
GCCCTGTTT ATTGCTTATG TTTGCTGGTT ATCGTGGTCA ACTATGTATC TGACACACCA 1260
TTATTTGTA GACCTTATGG CAGGTCTGT GCTGTCATAC GTTATTTCC AGTACACAAA 1320
GTACACACAT TTACCAATTG TAGATACATC TCTTTTTGC AGATGGTCAT AGACTTGAAT 1380
TGACAAATAC GATATATCAA ACAGTGATCC ATTGGCTGCA GATTCAAACG ATATCGAAAG 1440
TGTCCCTTG TCCAACCTTG AACATTGACTT TGATCTTAAT ATGACTGATG AACCCAGTGT 1500
AAGCCCTTCG TTATTTGATG GATCTACTTC TGTTCTCGT TCGTCCGCCA CGTCTATAAC 1560
GTCACTAGGT GTAAAGAGGG CTTAATGAGT ATTTTATCTG CAATTACGGA TACGGTTGGT 1620
CTTATGTAGA TACATATAAA TATATATCTT TTTCTTCTT TTTCTTAGTC AGGATTGTCG 1680
TTTACGATAA TATACATGTA GTTTATTTAA TCACATACCA CTGATTATCT TTAGAATTAA 1740
ATAAATTTT GAAATAAAATG GGTGGCTTT AATGGTGTCT ATGTTAAGTG AGGCTTTAG 1800
AATGCTCTTC CTGCTTGT TATTATATGT GTATGAAAGA TATGTATGTA TTTACATGTC 1860
TTTGTAGCGT CCCCAGTCAA AACCTGTGCG CTATACCTAA ATGGATTGAT AATCTTCATT 1920
CACTAATTCT AAAATAGACT TCTTCCCCAA AGAACGGTGT AACGATGAGG CTCTATCCAG 1980
CTCCTTATCT AAATCAACTT TAACGATGGA TGATCTTATG ACACGGGGAT CTTTCTTTAA 2040

AGTTCTTAGA ATTCAGACT GTACCGCAGC TGATGAATCA AACAGCATTA AAAAGTGATA 2100
TGCTCGAAAA TGTTCCTTCTT CATTATTTA GGAAGATACC TTATGCCAT 2160
GGCTACAATG TCCCTCACCA CACCTCTGTT TTGAATAATC AGTTCCCAGA TTGTGGAAGA 2220
CAATTCTTT GCTTCCAACCT TTGGCCATT GGAGTTGGTT ATGCGAACAA GTCCGATCAG 2280
CTCATAAAGC ATCTTAGTGA AAAGGCTGGT TTTGCGTTAT TCTTCCTCT GTTGAAGCTT 2340

SEQ ID NO : 8

SEQUENCE LENGTH : 401

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Ala Asn Pro Phe Ser Arg Trp Phe Leu Ser Glu Arg Pro Pro
1 5 10 15
Asn Cys His Val Ala Asp Leu Glu Thr Ser Leu Asp Pro His Gln
20 25 30
Thr Leu Leu Lys Val Gln Lys Tyr Lys Pro Ala Leu Ser Asp Trp
35 40 45
Val His Tyr Ile Phe Leu Gly Ser Ile Met Leu Phe Val Phe Ile
50 55 60
Thr Asn Pro Ala Pro Trp Ile Phe Lys Ile Leu Phe Tyr Cys Phe
65 70 75
Leu Gly Thr Leu Phe Ile Ile Pro Ala Thr Ser Gln Phe Phe Phe
80 85 90
Asn Ala Leu Pro Ile Leu Thr Trp Val Ala Leu Tyr Phe Thr Ser
95 100 105
Ser Tyr Phe Pro Asp Asp Arg Arg Pro Pro Ile Thr Val Lys Val
110 115 120

Leu Pro Ala Val Glu Thr Ile Leu Tyr Gly Asp Asn Leu Ser Asp
 125 130 135
 Ile Leu Ala Thr Ser Thr Asn Ser Phe Leu Asp Ile Leu Ala Trp
 140 145 150
 Leu Pro Tyr Gly Leu Phe His Phe Gly Ala Pro Phe Val Val Ala
 155 160 165
 Ala Ile Leu Phe Val Phe Gly Pro Pro Thr Val Leu Gln Gly Tyr
 170 175 180
 Ala Phe Ala Phe Gly Tyr Met Asn Leu Phe Gly Val Ile Met Gln
 185 190 195
 Asn Val Phe Pro Ala Ala Pro Pro Trp Tyr Lys Ile Leu Tyr Gly
 200 205 210
 Leu Gln Ser Ala Asn Tyr Asp Met His Gly Ser Pro Gly Gly Leu
 215 220 225
 Ala Arg Ile Asp Lys Leu Leu Gly Ile Asn Met Tyr Thr Thr Ala
 230 235 240
 Phe Ser Asn Ser Ser Val Ile Phe Gly Ala Phe Pro Ser Leu His
 245 250 255
 Ser Gly Cys Ala Thr Met Glu Ala Leu Phe Phe Cys Tyr Cys Phe
 260 265 270
 Pro Lys Leu Lys Pro Leu Phe Ile Ala Tyr Val Cys Trp Leu Trp
 275 280 285
 Trp Ser Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Met
 290 295 300
 Ala Gly Ser Val Leu Ser Tyr Val Ile Phe Gln Tyr Thr Lys Tyr
 305 310 315
 Thr His Leu Pro Ile Val Asp Thr Ser Leu Phe Cys Arg Trp Ser
 320 325 330
 Tyr Thr Ser Ile Glu Lys Tyr Asp Ile Ser Lys Ser Asp Pro Leu

335	340	345
Ala Ala Asp Ser Asn Asp Ile Glu Ser Val Pro Leu Ser Asn Leu		
350	355	360
Glu Leu Asp Phe Asp Leu Asn Met Thr Asp Glu Pro Ser Val Ser		
365	370	375
Pro Ser Leu Phe Asp Gly Ser Thr Ser Val Ser Arg Ser Ser Ala		
380	385	390
Thr Ser Ile Thr Ser Leu Gly Val Lys Arg Ala		
395	400	

SEQ ID NO : 9

SEQUENCE LENGTH : 5340

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION

AGCGCTTCTA TTTTCTCCC CACCGGGAGG CGAAATGGC ACATTTTTT TCTTTGCTT	60
CTGTGTTTT GCTGTAATT TTGGCATCTG CTATTGTATG AAGATAACGC GTGTTCCGT	120
GGAAATAGCC GGAAATTTG CCGGAATAT GACGGACATG ATTTAACACC CGTCGAAATG	180
AAAAAAGCCA AGGTAAGAAA GTGGCAATAT TTTTCTACA AATAGATCTG CTGTCCCTTA	240
CATGATTACC ATACATATAT ATATTTATTA CACACATCTG TCAGAGGTAG CTAGCGAAGG	300
TGTCACTGAA ATATTTTTG TTCCAGTTAG TATAAATACG GAGGTAGAAC AGCTCTCCGC	360
GTGTATATCT TTTTTGCGC TATACAAGAA CAGGAAGAAC GCATTTCCAT ACCTTTTCT	420
CCTTACAGGT GCCCTCTGAG TAGTGTACG AACGAGGAAA AAGATTAATA TTACTGTTT	480
TATATTCAA AAGAGTAAAG CCGTTGCTAT ATACGAATAT GACGATTACC GTGGGGGATG	540
CAGTTTCGGA GACGGAGCTG GAAAACAAA GTCAAAACGT GGTACTATCT CCCAAGGCCAT	600
CTGCTTCTTC AGACATAACC ACAGATGTTG ATAAAGACAC ATCGTCTTCT TGGGATGACA	660
AATCTTGCT GCCTACAGGT GAATATATTG TGGACAGAAA TAAGCCCCAA ACCTACTTGA	720

ATAGCGATGA TATCGAAAAA GTGACAGAAT CTGATATTT CCCTCAGAAA CCTCTGTTT 780
CATTCTTCCA CTCTAAGAAA ATTCCAGAAG TACCACAAAC CGATGACGAG AGGAAGATAT 840
ATCCTCTGTT CCATACAAAT ATTATCTCTA ACATGTTTT TTGCTGGTT CTACCCATCC 900
TGCAGTTGG TTATAAGAGA ACGATACAGC CGAACGATCT CTTCAAAATG GATCCGAGGA 960
TGTCTATAGA GACCCTTTAT GACCACTTG AAAAAAACAT GATTTACTAT TTTGAGAAGA 1020
CGAGGAAAAA ATACCGTAAA AGACATCCAG AAGCGACAGA AGAAGAGGTT ATGGAAAATG 1080
CCAAACTACC TAAACATACA GTTCTGAGAG CTTTATTATT CACTTTAAG AAACAGTACT 1140
TCATGTCGAT AGTGTGTTGCA ATTCTCGCTA ATTGTACATC CGGTTTAAC CCCATGATTA 1200
CCAAGAGGCT AATTGAGTTT GTCGAAGAAA AGGCTATTT TCATAGCATG CATGTTAAC 1260
AAGGTATTGG TTACGCTATT GGTCCATGTT TGATGATGTT CGTTAACGGG TTGACGTTCA 1320
ATCATTCTT TCATACATCC CAACTGACTG GTGTGCAAGC TAAGTCTATT CTTACTAAAG 1380
CTGCCATGAA GAAAATGTTT AATGCATCTA ATTATGCGAG ACATTGTTT CCTAACGGTA 1440
AAGTGACTTC TTTGTAACA ACAGATCTCG CTAGAATTGA ATTTGCCTTA TCTTTTCAGC 1500
CGTTTTGGC TGGGTTCCCT GCAATTGGC CTATTGCAAT TGTGTTATTG ATCGTTAAC 1560
TTGGAGCCAT TGCCTTAGTT GGGATTGGTA TTTTTTCGG TGGGTTTTTC ATATCCTTAT 1620
TTGCATTAA GTTAATTCTG GGCTTAGAA TTGCTGCGAA CATCTCACT GATGCTAGAG 1680
TTACCATGAT GAGAGAAGTG CTGAATAATA TAAAAATGAT TAAATATTAT ACGTGGGAGG 1740
ATGCGTATGA AAAAAATATT CAAGATATTA GGACCAAAGA GATTTCTAA GTTAGAAAAA 1800
TGCCTACTTC AAGAAATTTC TTGATTGCTA TGGCCATGTC TTTGCCTAGT ATTGCTTCAT 1860
TGGTCACTTT CCTTGCAATG TACAAAGTTA ATAAGGAGG CAGGCAACCT GGTAAATATT 1920
TTGCCCTCTTT ATCTTTATTG CAGGTCTTGA GTTTGCAAAT GTTTCTTA CCTATTGCTA 1980
TGGTACTGG AATTGACATG ATCATGGAT TGGGCCGTTT GCAAAGCTTA TTGGAGGCTC 2040
CAGAAGATGA TCCAAATCAG ATGATTGAAA TGAAGCCCTC TCCTGGCTTT GATCCAAAT 2100
TGGCTCTAAA AATGACACAT TGCTCATTG AGTGGGAAGA TTATGAATTA AACGACGCTA 2160
TTGAAGAAGC AAAAGGAGAA GCTAAAGATG AAGGTAAAAA GAACAAAAAA AAGCCTAAGG 2220
ATACATGGGG TAAGCCATCT GCAACTACTA ATAAGGCCAA AACATGGAC AATATGTTGA 2280
AAGACAGAGA CGGCCCCGAA GATTTAGAAA AAACCTCGTT TAGGGGTTTC AAGGACTTGA 2340
ACTTCGATAT TAAAAAGGGC GAATTATTAA TGATTACGGG ACCTATTGGT ACTGCTAAAT 2400
CTTCATTATT GAATGCGATG GCAGGATCAA TGAGAAAAAT TGATGCTAAG GTTGAAGTCA 2460

ACGGGGACTT ATTAATGTGT GGTTATCCAT GGATTCAAAA TGCATCTGTA AGAGATAACA 2520
TCATATTCCG TTCACCATTG AATAAAGAAA AGTATGATGA AGTAGTTCGT GTTTGCTCTT 2580
TGAAAGCTGA TCTGGATATT TTACCGGCAG GCGATATGAC CGAAATTGGG CAACGTGGTA 2640
TTACTTTATC TGGTGGTCAA AAGGCACGTA TCAATTAGC CAGGTCTGTT TATAAGAAGA 2700
AGGATATTG TGTATTCGAC GATGTCCTAA GTGCTGTCGA TTCTCGTGT AGTAAACACA 2760
TCATGGATGA ATGTCTAACCG GGAATGCTTG CTAATAAAAC CAGAATTGTA GCAACGCATC 2820
AGTTGTCACT GATTGAGAGA GCTTCTAGAG TCATCGTTT AGGTACTGAT GGCCAAGTCG 2880
ATATTGGTAC TGTTGATGAG CTAAAAGCTC GTAATCAAAC TTTGATAAAT CTTTTACAAT 2940
TCTCTCTCA AAATTCCGAG AAAGAGGATG AAGAACACGA AGCGGTGTT TCCGGTGAAT 3000
TGGGACAACG AAAATATGAA CCAGAGGTA AGGAATTGAC TGAACTGAAG AAAAAGGCTA 3060
CAGAAATGTC ACAAACTGCA AATAGTGGTA AAATTGTAAC GGATGGTCAT ACTAGTAGTA 3120
AAGAAGAAAG ACCAGTCAT AGTATCAGTC TGAAAATATA CCGTGAATAC ATTAAGCTG 3180
CACTAGGTAA GTGGGGTTTT ATCGCACTAC CGTTGTATGC AATTTAGTC GTTGGAAACCA 3240
CATTCTGCTC ACTTTTTCT TCCGTTGGT TATCTTACTG GACTGAGAAT AAAATTCAAAA 3300
ACAGACCACC CAGTTTTAT ATGGGTCTT ACTCCTTCTT TGTGTTGCT GCTTCATAT 3360
TCATGAATGG CCAGTTCACCC ATACTTGCG CAATGGGTAT TATGGCATCG AAATGGTTAA 3420
ATTTGAGGGC TGTGAAAAGA ATTTACACA CTCCAATGTC ATACATAGAT ACCACACCTT 3480
TGGGACCTAT TCTGAACAGA TTCACAAAAG ATACAGATAG CTTAGATAAT GAGTTAACCG 3540
AAAGTTTACG GTTGATGACA TCTCAATTG CTAATATTGT AGGTGTTGC GTCATGTGTA 3600
TTGTTTACTT GCCGTGGTTT GCTATCGCAA TTCCGTTCT TTTGGTCATC TTTGTTCTGA 3660
TTGCTGATCA TTATCAGAGT TCTGCTAGAG AAATTAAAAG ACTTGAAGCT GTGCAACGGT 3720
CTTTTGTAA CAATAATTAA AATGAAGTTT TGGGTGGGAT GGATACAATC AAAGCATAACC 3780
GAAACTCAGGA ACGATTTTG GCGAAATCAG ATTTTTGAT CAACAAGATG AATGAGGCCGG 3840
GATACCTTGT AGTTGTCCTG CAAAGATGGG TAGGTATTTT CCTTGATATG GTTGCTATCG 3900
CATTTGCACT AATTATTACG TTATTGTGTC TTACGAGAGC CTTTCCTATT TCCCGGGCTT 3960
CAGTTGGTGT TTTGTTGACT TATGTATTAC AATTGCTGG TCTATTAAAT ACCATTTAA 4020
GGGCAATGAC TCAAACAGAG AATGACATGA ATAGTGCCGA AAGATTGGTA ACATATGCAA 4080
CTGAACTACC ACTAGAGGCA TCCTATAGAA AGCCCCAAAT GACACCTCCA GAGTCATGGC 4140
CCTCAATGGG CGAAATAATT TTTGAAAATG TTGATTTGC CTATAGACCT GGTTTACCTA 4200

TAGTTTAAA AAATCTTAAC TTGAATATCA AGAGTGGGA AAAAATTGGT ATCTGTGGTC 4260
GTACAGGTGC TCGTAAGTCC ACTATTATGA GTGCCCTTA CAGGTGAAT GAATTGACCG 4320
CAGGTAAAAT TTTAATTGAC AATGTGATA TAAGTCAGCT GGGACTTTTC GATTTAAGAA 4380
GAAAATTAGC CATCATTCCA CAAGATCCAG TATTATTTAG GGGTACGATT CGCAAGAACT 4440
TAGATCCATT TAATGAGCGT ACAGATGACG AATTATGGGA TGCATTGGTG AGAGGTGGTG 4500
CTATGCCAA GGATGACTTG CCGGAAGTGA ATTGCAAAA ACCTGATGAA AATGGTACTC 4560
ATGGTAAAAT GCATAAGTTC CATTAGATC AAGCAGTGGA AGAAGAGGGC TCCAATTCT 4620
CCTTAGGTGA GAGACAACTA TTAGCATTAA CAAGGGCATT GGTCCGCCAA TCAAAAAAT 4680
TGATTTGGG TGAGGCTACA TCCTCACTGG ACTACGAAAC GGATGGCAAATCCTAACAC 4740
CTATGTTGA GGAATTGGA GATTGTACAA TTTTGTGTAT TCCTCACAGA CTGAAGACCA 4800
TTGTAATTG TGATCGTATT CTTGTTTGTAG AGAAGGGTGA AGTCCGAGAA TCCGATACAC 4860
CATGGACGTT GTTGTGCTAA GAAGATACTA TTTTCAGAAG CATGTGTTCT AGATCTGGTA 4920
TTGTCGAAAA TGATTTCGAG AACAGAAAGTT AATTTATATT ATTTGTGTGA TGATTTTC 4980
CTTTTATTGTT TTTATATGTT GCCGATGGTA CAAATTAGTA CTAGAAAAGA AAACCCACTA 5040
CTATGACTTG CAGAAAAAGT TATGTGTGCC ATAGATAGAT ATAATTGCAAT ACCCACATCG 5100
TATACTCAA ATTCCGAAAA GAACATTCA TTTTTTATGA GGCAAACGTA AGAACGCTTC 5160
GGTCCTTTT TCATTCAGA AATATATATT TATACATCAT TTTCAGAAGA TATTCAAAGA 5220
ACTTATTGGG ATGTCTATT ACTGAATAAA GTATACACAA AAAACGAATT TAAATGGAA 5280
GGCATAAATA GAAAACCTAG AACTGAAAAT CCTAAAACCG AAGGATATTG CAAATACCTA 5340

SEQ ID NO: 10

SEQUENCE LENGTH: 1477

SEQUENCE TYPE: amino acid

STRANDEDNESS: Single

TOPOLOGY: linear

MOLECULE TYPE: peptide

SEQUENCE DESCRIPTION:

Met Thr Ile Thr Val Gly Asp Ala Val Ser Glu Thr Glu Leu Glu

Asn Lys Ser Gln Asn Val Val Leu Ser Pro Lys Ala Ser Ala Ser
20 25 30
Ser Asp Ile Ser Thr Asp Val Asp Lys Asp Thr Ser Ser Ser Trp
35 40 45
Asp Asp Lys Ser Leu Leu Pro Thr Gly Glu Tyr Ile Val Asp Arg
50 55 60
Asn Lys Pro Gln Thr Tyr Leu Asn Ser Asp Asp Ile Glu Lys Val
65 70 75
Thr Glu Ser Asp Ile Phe Pro Gln Lys Arg Leu Phe Ser Phe Leu
80 85 90
His Ser Lys Lys Ile Pro Glu Val Pro Gln Thr Asp Asp Glu Arg
95 100 105
Lys Ile Tyr Pro Leu Phe His Thr Asn Ile Ile Ser Asn Met Phe
110 115 120
Phe Trp Trp Val Leu Pro Ile Leu Arg Val Gly Tyr Lys Arg Thr
125 130 135
Ile Gln Pro Asn Asp Leu Phe Lys Met Asp Pro Arg Met Ser Ile
140 145 150
Glu Thr Leu Tyr Asp Asp Phe Glu Lys Asn Met Ile Tyr Tyr Phe
155 160 165
Glu Lys Thr Arg Lys Lys Tyr Arg Lys Arg His Pro Glu Ala Thr
170 175 180
Glu Glu Glu Val Met Glu Asn Ala Lys Leu Pro Lys His Thr Val
185 190 195
Leu Arg Ala Leu Leu Phe Thr Phe Lys Lys Gln Tyr Phe Met Ser
200 205 210
Ile Val Phe Ala Ile Leu Ala Asn Cys Thr Ser Gly Phe Asn Pro
215 220 225
Met Ile Thr Lys Arg Leu Ile Glu Phe Val Glu Glu Lys Ala Ile

230	235	240
Phe His Ser Met His Val Asn Lys Gly Ile Gly Tyr Ala Ile Gly		
245	250	255
Ala Cys Leu Met Met Phe Val Asn Gly Leu Thr Phe Asn His Phe		
260	265	270
Phe His Thr Ser Gln Leu Thr Gly Val Gln Ala Lys Ser Ile Leu		
275	280	285
Thr Lys Ala Ala Met Lys Lys Met Phe Asn Ala Ser Asn Tyr Ala		
290	295	300
Arg His Cys Phe Pro Asn Gly Lys Val Thr Ser Phe Val Thr Thr		
305	310	315
Asp Leu Ala Arg Ile Glu Phe Ala Leu Ser Phe Gln Pro Phe Leu		
320	325	330
Ala Gly Phe Pro Ala Ile Leu Ala Ile Cys Ile Val Leu Leu Ile		
335	340	345
Val Asn Leu Gly Pro Ile Ala Leu Val Gly Ile Gly Ile Phe Phe		
350	355	360
Gly Gly Phe Phe Ile Ser Leu Phe Ala Phe Lys Leu Ile Leu Gly		
365	370	375
Phe Arg Ile Ala Ala Asn Ile Phe Thr Asp Ala Arg Val Thr Met		
380	385	390
Met Arg Glu Val Leu Asn Asn Ile Lys Met Ile Lys Tyr Tyr Thr		
395	400	405
Trp Glu Asp Ala Tyr Glu Lys Asn Ile Gln Asp Ile Arg Thr Lys		
410	415	420
Glu Ile Ser Lys Val Arg Lys Met Gln Leu Ser Arg Asn Phe Leu		
425	430	435
Ile Ala Met Ala Met Ser Leu Pro Ser Ile Ala Ser Leu Val Thr		
440	445	450

Phe Leu Ala Met Tyr Lys Val Asn Lys Gly Gly Arg Gln Pro Gly
455 460 465
Asn Ile Phe Ala Ser Leu Ser Leu Phe Gln Val Leu Ser Leu Gln
470 475 480
Met Phe Phe Leu Pro Ile Ala Ile Gly Thr Gly Ile Asp Met Ile
485 490 495
Ile Gly Leu Gly Arg Leu Gln Ser Leu Leu Glu Ala Pro Glu Asp
500 505 510
Asp Pro Asn Gln Met Ile Glu Met Lys Pro Ser Pro Gly Phe Asp
515 520 525
Pro Lys Leu Ala Leu Lys Met Thr His Cys Ser Phe Glu Trp Glu
530 535 540
Asp Tyr Glu Leu Asn Asp Ala Ile Glu Glu Ala Lys Gly Glu Ala
545 550 555
Lys Asp Glu Gly Lys Lys Asn Lys Lys Arg Lys Asp Thr Trp
560 565 570
Gly Lys Pro Ser Ala Ser Thr Asn Lys Ala Lys Arg Leu Asp Asn
575 580 585
Met Leu Lys Asp Arg Asp Gly Pro Glu Asp Leu Glu Lys Thr Ser
590 595 600
Phe Arg Gly Phe Lys Asp Leu Asn Phe Asp Ile Lys Lys Gly Glu
605 610 615
Phe Ile Met Ile Thr Gly Pro Ile Gly Thr Gly Lys Ser Ser Leu
620 625 630
Leu Asn Ala Met Ala Gly Ser Met Arg Lys Ile Asp Gly Lys Val
635 640 645
Glu Val Asn Gly Asp Leu Leu Met Cys Gly Tyr Pro Trp Ile Gln
650 655 660
Asn Ala Ser Val Arg Asp Asn Ile Ile Phe Gly Ser Pro Phe Asn

665 670 675
Lys Glu Lys Tyr Asp Glu Val Val Arg Val Cys Ser Leu Lys Ala
680 685 690
Asp Leu Asp Ile Leu Pro Ala Gly Asp Met Thr Glu Ile Gly Glu
695 700 705
Arg Gly Ile Thr Leu Ser Gly Gly Gln Lys Ala Arg Ile Asn Leu
, 710 715 720
Ala Arg Ser Val Tyr Lys Lys Asp Ile Tyr Val Phe Asp Asp
725 730 735
Val Leu Ser Ala Val Asp Ser Arg Val Gly Lys His Ile Met Asp
740 745 750
Glu Cys Leu Thr Gly Met Leu Ala Asn Lys Thr Arg Ile Leu Ala
755 760 765
Thr His Gln Leu Ser Leu Ile Glu Arg Ala Ser Arg Val Ile Val
770 775 780
Leu Gly Thr Asp Gly Gln Val Asp Ile Gly Thr Val Asp Glu Leu
785 790 795
Lys Ala Arg Asn Gln Thr Leu Ile Asn Leu Leu Gln Phe Ser Ser
800 805 810
Gln Asn Ser Glu Lys Glu Asp Glu Glu Gln Glu Ala Val Val Ser
815 820 825
Gly Glu Leu Gly Gln Leu Lys Tyr Glu Pro Glu Val Lys Glu Leu
830 835 840
Thr Glu Leu Lys Lys Ala Thr Glu Met Ser Gln Thr Ala Asn
845 850 855
Ser Gly Lys Ile Val Ala Asp Gly His Thr Ser Ser Lys Glu Glu
860 865 870
Arg Ala Val Asn Ser Ile Ser Leu Lys Ile Tyr Arg Glu Tyr Ile
875 880 885

Lys Ala Ala Val Gly Lys Trp Gly Phe Ile Ala Leu Pro Leu Tyr
890 895 900
Ala Ile Leu Val Val Gly Thr Thr Phe Cys Ser Leu Phe Ser Ser
905 910 915
Val Trp Leu Ser Tyr Trp Thr Glu Asn Lys Phe Lys Asn Arg Pro
920 925 930
Pro Ser Phe Tyr Met Gly Leu Tyr Ser Phe Phe Val Phe Ala Ala
935 940 945
Phe Ile Phe Met Asn Gly Gin Phe Thr Ile Leu Cys Ala Met Gly
950 955 960
Ile Met Ala Ser Lys Trp Leu Asn Leu Arg Ala Val Lys Arg Ile
965 970 975
Leu His Thr Pro Met Ser Tyr Ile Asp Thr Thr Pro Leu Gly Arg
980 985 990
Ile Leu Asn Arg Phe Thr Lys Asp Thr Asp Ser Leu Asp Asn Glu
995 1000 1005
Leu Thr Glu Ser Leu Arg Leu Met Thr Ser Gin Phe Ala Asn Ile
1010 1015 1020
Val Gly Val Cys Val Met Cys Ile Val Tyr Leu Pro Trp Phe Ala
1025 1030 1035
Ile Ala Ile Pro Phe Leu Leu Val Ile Phe Val Leu Ile Ala Asp
1040 1045 1050
His Tyr Gin Ser Ser Gly Arg Glu Ile Lys Arg Leu Glu Ala Val
1055 1060 1065
Gin Arg Ser Phe Val Tyr Asn Asn Leu Asn Glu Val Leu Gly Gly
1070 1075 1080
Met Asp Thr Ile Lys Ala Tyr Arg Ser Gin Glu Arg Phe Leu Ala
1085 1090 1095
Lys Ser Asp Phe Leu Ile Asn Lys Met Asn Glu Ala Gly Tyr Leu

1100 1105 1110
Val Val Val Leu Gin Arg Trp Val Gly Ile Phe Leu Asp Met Val
1115 1120 1125
Ala Ile Ala Phe Ala Leu Ile Ile Thr Leu Leu Cys Val Thr Arg
1130 1135 1140
Ala Phe Pro Ile Ser Ala Ala Ser Val Gly Val Leu Leu Thr Tyr
1145 1150 1155
Val Leu Gin Leu Pro Gly Leu Leu Asn Thr Ile Leu Arg Ala Met
1160 1165 1170
Thr Gin Thr Glu Asn Asp Met Asn Ser Ala Glu Arg Leu Val Thr
1175 1180 1185
Tyr Ala Thr Glu Leu Pro Leu Glu Ala Ser Tyr Arg Lys Pro Glu
1190 1195 1200
Met Thr Pro Pro Glu Ser Trp Pro Ser Met Gly Glu Ile Ile Phe
1205 1210 1215
Glu Asn Val Asp Phe Ala Tyr Arg Pro Gly Leu Pro Ile Val Leu
1220 1225 1230
Lys Asn Leu Asn Leu Asn Ile Lys Ser Gly Glu Lys Ile Gly Ile
1235 1240 1245
Cys Gly Arg Thr Gly Ala Gly Lys Ser Thr Ile Met Ser Ala Leu
1250 1255 1260
Tyr Arg Leu Asn Glu Leu Thr Ala Gly Lys Ile Leu Ile Asp Asn
1265 1270 1275
Val Asp Ile Ser Gin Leu Gly Leu Phe Asp Leu Arg Arg Lys Leu
1280 1285 1290
Ala Ile Ile Pro Gin Asp Pro Val Leu Phe Arg Gly Thr Ile Arg
1295 1300 1305
Lys Asn Leu Asp Pro Phe Asn Glu Arg Thr Asp Asp Glu Leu Trp
1310 1315 1320

Asp Ala Leu Val Arg Gly Gly Ala Ile Ala Lys Asp Asp Leu Pro
1325 1330 1335
Glu Val Lys Leu Gln Lys Pro Asp Glu Asn Gly Thr His Gly Lys
1340 1345 1350
Met His Lys Phe His Leu Asp Gln Ala Val Glu Glu Gly Ser
1355 1360 1365
Asn Phe Ser Leu Gly Glu Arg Gln Leu Leu Ala Leu Thr Arg Ala
1370 1375 1380
Leu Val Arg Gln Ser Lys Ile Leu Ile Leu Asp Glu Ala Thr Ser
1385 1390 1395
Ser Val Asp Tyr Glu Thr Asp Gly Lys Ile Gln Thr Arg Ile Val
1400 1405 1410
Glu Glu Phe Gly Asp Cys Thr Ile Leu Cys Ile Ala His Arg Leu
1415 1420 1425
Lys Thr Ile Val Asn Tyr Asp Arg Ile Leu Val Leu Glu Lys Gly
1430 1435 1440
Glu Val Ala Glu Phe Asp Thr Pro Trp Thr Leu Phe Ser Gln Glu
1445 1450 1455
Asp Ser Ile Phe Arg Ser Met Cys Ser Arg Ser Gly Ile Val Glu
1460 1465 1470
Asn Asp Phe Glu Asn Arg Ser
1475

SEQ ID NO : 11

SEQUENCE LENGTH : 26

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : Other nucleic acid(synthetic DNA)

SEQUENCE DESCRIPTION :

TTTGGTTAYA TGAAYYTNTT YGGNGT 26

SEQ ID NO : 12

SEQUENCE LENGTH : 29

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : Other nucleic acid(synthetic DNA)

SEQUENCE DESCRIPTION :

TCTACAAART ARTGGTGNGT NARRTACAT 29

SEQ ID NO : 13

SEQUENCE LENGTH : 2274

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION :

TTATATATAT TATTGATTG TTCCCTGTTGT TATTTAGTTT AGAACATCAGAC GACTACACCA 60
GAACCACAAT TCAACCAACA CTTATATAGA ACCTGGCTTG GAAAAAAAGTA ACATTTATCA 120
TTCCCTATACT TTTTTAGCAA ACATAATCCG TGTTTTACAT ATATTATTCA CCCAATATCA 180
TAACAAAAAAC AAACTGAATA ATGCCGTCTT CTATTTGCG TTCCAAAATA ATACAAAAAC 240
CGTACCAATT ATTCCACTAC TATTTCTTC TGGAGAAGGC TCCTGGTTCT ACAGTTAGTG 300
ATTGAATTT TGATACAAAC ATACAAACGA GTTTACGTAATTAAGCAT CATCATTGGA 360
CGGTGGGAGA AATATTCCAT TATGGGTTTT TGGTTTCCAT ACTTTTTTC GTGTTGTGG 420
TTTCCCAGC TTCATTTTT ATAAAATTAC CAATAATCTT AGCATTGCT ACTTGTTTT 480
TAATACCCCTT AACATCACAA TTTTTCTTC CTGCCTTGCC CGTTTCACT TGGTTGGCAT 540
TATATTTAC GTGTGCTAAA ATACCTCAAG AATGGAAACC AGCTATCACA GTTAAAGTTT 600
TACCAAGCTAT GGAAACAATT TTGTACGGCG ATAATTATC AAATGTTTG GCAACCATCA 660

CTACCGGAGT GTTAGATATA TTGCATGGT TACCATATGG GATTATTCAT TTCAGTTCC 720
CATTCTACT TGCTGCTATT ATATTTTAT TTGGGCCACC GACGGCATT AAGTCATTG 780
GATTGCCTT TGGTTATATG AACTGCTTG GAGTCTTGAT TCAAATGCCA TTCCCAGCTG 840
CTCCTCCATG GTACAAAAAC TTGCACGGAT TAGAACCAAGC TAATTATTCA ATGCACGGGT 900
CTCCTGGTGG ACTTGGAAAGG ATAGATAAAT TGTTAGGTGT TGATATGTAT ACCACAGGGT 960
TTTCCAATTG ATCAATCATT TTTGGGCCAT TCCCATCGTT ACATTCAAGGA TGGTGTATCA 1020
TGGAAAGTGTGTTTGTGTTTC CACGATTCAA GTTTGTGTGG GTTACATACG 1080
CATCTTGGCT TTGGTGGAGC ACCATGTATT TGACCCATCA CTACTTGTGTC GATTTGATTC 1140
GTGGAGCCAT GCTATCTTG ACTGTTTTG AGTTCACCAA ATATAAATAT TTGCCAAAAA 1200
ACAAGAAGG CCTTTCTGT CGTTGGTCAT ACACGTAAAT TGAAAAAATC GATATCCAAG 1260
AGATTGACCC TTTATCATAA ATTATATCC CTGTCAACAG CAATGATAAT GAAAGCAGAT 1320
TGTATACGAG AGTGTACCAA GAGTCTCAGG TTAGTCCCCC ACAGAGAGCT GAAACACCTG 1380
AAGCATTG AATGTCAAAT TTTCTAGGT CTAGACAAAG CTCAAAGACT CAGGTTCCAT 1440
TGAGTAATCT TACTAACAAAT GATCAAGTGT CTGGAATTAA CGAAGAGGAT GAAGAAGAAG 1500
AAGGCGATGA AATTTCATCG AGTACTCCTT CGGTGTTGA AGACGAACCA CAGGGTAGCA 1560
CATATGCTGC ATCCTCAGCT ACATCACTAG ATGATTTGGA TTCCAAAAGA AATTAGTAAA 1620
ATAACAGTTT CTATTAATTTCCTTCC TCCATTAA TGATTTATG CTCATAACCT 1680
ACACTATCTG TTTTAATTTCCTTCC TTTTATTATT GTTGAGTTCA TTTGCTGTTG 1740
ATTGAATATT TACAATTTG CATTAAATTAC CATCAATATA GAATGGCAC AGTTTTTTA 1800
AGTTTTTG TTTTGTGT TGTCTTCTT TTTTACATT AATGTGTTG GATTGTTTA 1860
GGTTCCCTTA TCCCTTAAGCC CCCTCAGAAT ACTATTTAT CTAATTAAATT TGTTTTATT 1920
TTCTGATATT TACCAATTGC TTTTCTTTT CGATATTTAT AATAGCATCC CCTAATAATT 1980
AATATACAAC TGTTCATAT ATATACGTGT ATGTCTGTGA GTGGTGGAAA CTGGACTCAA 2040
CATTCTATT AATGTGTACA AGAAAGCAGT GTTAATGCTA CTATTATAAT TTTGAGGTG 2100
CAAATCAAGA GGTTGGCAGC TTTCTTATGG CTATGACCGT GAATGAAGGC TTGTAAACCA 2160
CGTAATAAAC AAAAGCCAAC AAGTTTTT AGAGCCTTA ACAACATACG CAATGAGACT 2220
GATTGCAATA CTACAAGATA TAGCCAAAAA AATTGAATGC ATTTCAACAA CAAC 2274

SEQ ID NO : 14

SEQUENCE LENGTH : 471

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Ala Ser Ser Ile Leu Arg Ser Lys Ile Ile Gln Lys Pro Tyr
5 10 15
Gln Leu Phe His Tyr Tyr Phe Leu Ser Glu Lys Ala Pro Gly Ser
20 25 30
Thr Val Ser Asp Leu Asn Phe Asp Thr Asn Ile Gln Thr Ser Leu
35 40 45
Arg Lys Leu Lys His His His Trp Thr Val Gly Glu Ile Phe His
50 55 60
Tyr Gly Phe Leu Val Ser Ile Leu Phe Phe Val Phe Val Val Phe
65 70 75
Pro Ala Ser Phe Phe Ile Lys Leu Pro Ile Ile Leu Ala Phe Ala
80 85 90
Thr Cys Phe Leu Ile Pro Leu Thr Ser Gln Phe Phe Leu Pro Ala
95 100 105
Leu Pro Val Phe Thr Trp Leu Ala Leu Tyr Phe Thr Cys Ala Lys
110 115 120
Ile Pro Gln Glu Trp Lys Pro Ala Ile Thr Val Lys Val Leu Pro
125 130 135
Ala Met Glu Thr Ile Leu Tyr Gly Asp Asn Leu Ser Asn Val Leu
140 145 150
Ala Thr Ile Thr Thr Gly Val Leu Asp Ile Leu Ala Trp Leu Pro
155 160 165
Tyr Gly Ile Ile His Phe Ser Phe Pro Phe Val Leu Ala Ala Ile

170	175	180
Ile Phe Leu Phe Gly Pro Pro Thr Ala Leu Arg Ser Phe Gly Phe		
185	190	195
Ala Phe Gly Tyr Met Asn Leu Leu Gly Val Leu Ile Gln Met Ala		
200	205	210
Phe Pro Ala Ala Pro Pro Trp Tyr Lys Asn Leu His Gly Leu Glu		
215	220	225
Pro Ala Asn Tyr Ser Met His Gly Ser Pro Gly Gly Leu Gly Arg		
230	235	240
Ile Asp Lys Leu Leu Gly Val Asp Met Tyr Thr Thr Gly Phe Ser		
245	250	255
Asn Ser Ser Ile Ile Phe Gly Ala Phe Pro Ser Leu His Ser Gly		
260	265	270
Cys Cys Ile Met Glu Val Leu Phe Leu Cys Trp Leu Phe Pro Arg		
275	280	285
Phe Lys Phe Val Trp Val Thr Tyr Ala Ser Trp Leu Trp Trp Ser		
290	295	300
Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Ile Gly Gly		
305	310	315
Ala Met Leu Ser Leu Thr Val Phe Glu Phe Thr Lys Tyr Lys Tyr		
320	325	330
Leu Pro Lys Asn Lys Glu Gly Leu Phe Cys Arg Trp Ser Tyr Thr		
335	340	345
Glu Ile Glu Lys Ile Asp Ile Gln Glu Ile Asp Pro Leu Ser Tyr		
350	355	360
Asn Tyr Ile Pro Val Asn Ser Asn Asp Asn Glu Ser Arg Leu Tyr		
365	370	375
Thr Arg Val Tyr Gln Glu Ser Gln Val Ser Pro Pro Gln Arg Ala		
380	385	390

Glu Thr Pro Glu Ala Phe Glu Met Ser Asn Phe Ser Arg Ser Arg

395 400 405

Gln Ser Ser Lys Thr Gln Val Pro Leu Ser Asn Leu Thr Asn Asn

410 415 420

Asp Gln Val Ser Gly Ile Asn Glu Glu Asp Gln Glu Glu Gly

425 430 435

Asp Glu Ile Ser Ser Ser Thr Pro Ser Val Phe Glu Asp Glu Pro

440 445 450

Gln Gln Ser Thr Tyr Ala Ala Ser Ser Ala Thr Ser Val Asp Asp

455 460 465

Leu Asp Ser Lys Arg Asn

470

SEQ ID NO : 15

SEQUENCE LENGTH : 243

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

SEQUENCE DESCRIPTION

TTTGAATAAT TTGAAATTAA AAATTAATCC AATGGAAAAA ATTGGTATTT GTGGAAGAAC 60

CGGTGCTGGT AAATCATCAA TTATGACAGC ATTATATCGA TTATCAGAAAT TAGAACTGGG 120

GAAAATTATT ATTGATGATA TTGATATTTC AACTTTGGGT TTAAAAGATC TTGATCAA 180

ATTATGAATT ATTCCCTCAAG ATCCAGTATT ATTCCGAGGT TCAATTCCGA AAAACTTGGG 240

TCC 243

SEQ ID NO : 16

SEQUENCE LENGTH : 80

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Leu Lys Asn Leu Asn Phe Lys Ile Asn Pro Met Glu Lys Ile Gly

5

10

15

Ile Cys Gly Arg Thr Gly Ala Gly Lys Ser Ser Ile Met Thr Ala

20

25

30

Leu Tyr Arg Leu Ser Glu Leu Glu Leu Gly Lys Ile Ile Ile Asp

35

40

45

Asp Ile Asp Ile Ser Thr Leu Gly Leu Lys Asp Leu Arg Ser Lys

50

55

60

Leu Ser Ile Ile Pro Cln Asp Pro Val Leu Phe Arg Gly Ser Ile

65

70

75

Arg Lys Asn Leu Asp

80

SEQ ID NO : 17

SEQUENCE LENGTH : 1601

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : genomic DNA

ANTI-SENSE : Yes

SEQUENCE DESCRIPTION :

AGGAAGATCA CTTGCATCAA AGATGGAGGA AGTGGTACTG GCAGGACGAT CAATCAAATC 60

AGCACCGAGGA CTAGGTAACG GCTCAGGTGA TGATCAACCC ACGGACCATT CATGATCGGT 120

GTTAGCAAGT TCCATATTGT TAAGACCACT CATGAAGGCT ACTGCATTAG GGTTTGAGT 180

AAAAGAATCC CTTCCAAGTA AGTATGGGCT GCCGGTACGA GCCAAGGAGT TGCTGGTTT 240

TTCGGAAAGA CCATGACCGT GGATAACAAA CTCGTATTCC CAACGAAGGA TTTTACCACT 300

TTGCAACTGT GGGAGGGCTA GCTTTGAGC AAAAACGAAG CATATAATAG CTAACACAT 360

ACCGCCGACC	AAATCTACAA	AGTAGTGGTG	GGTAAGGTAC	ATAGTACACC	AGCAAAGCCA	420
TAGAACATAT	CCATAAAAGC	AGAAGCGGT	TCGAGGAAAC	ACATGCGAAA	GGAAAAGTGC	480
TTCCAGCATG	GCCCATCCAG	CGTGAAGAGA	TGGAAAGGCA	CCAAAAACAA	CCGGAGAGTT	540
AGAAAAACCA	TCAGTGTAAA	TGCTAGTGCC	GAAGAGAGCA	TCAATACGGG	CCAATCCACC	600
AGGAGAGCCA	CGTACTGCAT	ACGTGGCAGG	TTCTAAACCA	TACATATTTC	CATACCAAGG	660
AGGAGAACAG	GGGAAAGCCA	TTTGGATAAG	AACACCAAAT	AAATTCAATAT	AACCAAAAGT	720
TCGAGCCCCAA	ACTGGAAGAG	TTCCAGGAGG	TGCAAACATG	AAAACAATAA	ATGAAATGAT	780
AAAAGGAGCC	GAATAATGCA	TGACTCCATA	TGGAACCCAG	GCCAAAATAT	CAAGGATGCT	840
ATCCGTGGTT	TTCGAGAGAA	GACTAGAAAC	ATTAGAGCCA	TAAGAATAT	TTTCAAGTGT	900
GGGTAAAACA	CGAACCCATA	TGGGTGGACG	CCAGCGTTCT	GGAATAAAACC	TACAAGACTA	960
AAATAAAATT	GCCCCAGGTGA	TGATAACAAT	GGCAGGAAAAA	AAAATTGCG	GTGTTAAAGG	1020
AACGGTCAAC	GCAATGGCCA	AAAGACAGGC	AATGCCAAAT	TTCCCCCAGA	ATCCACGGAGA	1080
TTCAATGACA	ATACAAGCAA	AAATCAAATT	ACCTGCTAGA	AACACATATT	GCAAATGTGT	1140
CCATGACCAT	TTCGTATTGC	GTAGCAAACG	AAATGTAGGC	ATAGGGTTA	ACCTTGTTC	1200
CAACITGTAT	TGGGATGCTC	GGTTACACCC	AGCAAGGCC	TTTTTAAGG	TCGAAAGAGC	1260
AGACATTGCT	TCAAAGAATT	ATCAGACTAA	AAAAGGAAG	CGTACGAAAAA	AAATTGCGTA	1320
ACCAATTAAAC	CGGAAAACCA	AAAGGAAAAAA	AACGAATTTC	TATGAAGGAA	AGAAAGTAGGC	1380
TATTAATGCA	AAGTGTCAAG	CACTAAAAG	TAGGGATGTA	AAATATTAA	AAAAAGATGG	1440
ACCGATTAAAC	CAATGTCAG	CTCACAGTTG	CCAGCAATCA	GGGCTATTTT	TTTATTTTT	1500
TTATAAAATT	GCTAATTATA	TATAATATAA	TTAGTTTATT	AACTTGCTTT	TCCTCAAAAA	1560
ACCAATTGCA	CAAAGGAAC	TTTGCAGAGG	CAAAAAAGCT	T		1601

SEQ_ID NO : 18

SEQUENCE LENGTH : 1601

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : mRNA

ANTI-SENSE : Yes

SEQUENCE DESCRIPTION :

AGGAAGAUGA CUUGCAUCAA AGAUGGAGGA AGUGGUACUG GCAGGACGAU CAAUCAAAUC 60
AGCAGCAGGA CUAGGUACG GCUCAGGUCA UGAUGAACCC ACCGACCAUU CAUGAUCCGU 120
GUUAGCAAGU UCCAUAUUCU UAAGACCACU CAUGAAGGCU ACUGCAUUAG GGUUUUUGAGU 180
AAAAGAAUCC CUUCCAAGUA AGUAUGGGCU GCCGGUACGA GCCAAGGAGU UGCUGGUUUU 240
UUCGGAAAGA CCAUGACCGU GGAAUACAAA CUCCGUUUCU CAACGAAGGA UUUUACCAU 300
UUGCAACUGU GGGAGGCCUA GCUUUUGAGC AAAAACGAAG CAUAAUUAAG CUAAACACAU 360
ACCGCCGACC AAAUCUACAA AGUAGUGGUG GGUAAGGUAC AUAGUACACC AGCAAAGGCCA 420
UAGAACAUAU CCAUAAAAGC AGAAGCGGU ACGAGGAAAC ACAUGCAGA ..24AAAGUGC 480
UUCAGCAUG CCCAUCCAG CGUGAAGAGA UGGAAAGGCA CCAA ..ACAA CGGGAGAGGU 540
AGAAAAACCA UCAGUGUAAA UCCUAGUGCC GAAGAGAGCA UCAAUACGCC CCAAUCCACC 600
ACCGAGCCCAC GUACUGCAU ACCUGGCCAGG UUCUAAACCA UACAUUUU CAUACCAACG 660
AGGAGAACAG CCCAAAGCCA UUUGGAUAAG AACACCAAAU AAAUUCAUU AACCAAACU 720
UCCAGCCCCA ACUGGAAGAG UUCCAGGAGG UGCAAAGAUG AAAAGAAUAA AUGAAUUGAU 780
AAAAGGAGCC GAAUAAUGCA UGACUCCAU UGGAACCCAG GCCAAAUAU CAAGGAUGCU 840
AUGCGUGGUU UUCGAGAGAA GACUAGAAAG AUUAGGCCA UAAAGAAUAU UUCAAGUGU 900
CGGUAAAACA CGAACCCAU UGGGUGGACG CCAGGGUUCU GGAAUAAACC UACAAGAGUA 960
AAAUAUUU GCCCAGGUCA UGAUAACAAU GGCAAGAAAA AAAAUUUGGC GUGUUAAGG 1020
AACGGUCAAC GCAAUGGCCA AAAGACAGGC AAUGCCAAU UUCCCCAGA AUCCAGGACA 1080
UUCAAUGACA AUACAAGCAA AAAUCAAAUU ACCUGGUAGA AACACAUUU GCAAUUGGU 1140
CCAUGACCAU UUCGUAUUGC GUAGCAAACG AAAUGUAGGC AUAGGGUUUA AGCUUGGUUC 1200
CAACUUGUAU UGGGAUGGCUC CGUACACGC ACCAAGGCCG UUUUUUAAGG UCGAAAGAGC 1260
AGACAUUGCU UCAAAGAAUU AUCAGAGUA AAAAGGGAAG CGUACGAAAA AAAUUUCCUA 1320
AGGAUUUAAC CGGAAAACUA AAGGAAAAAA AAGGAAUUUUA UAGAAGGAA ACAAAAGUAGC 1380
UAUAAAUGC AACUGUCAAG CACUAAAAG UAGCGAUGUA AAAUAAUUA AAAAGAUGG 1440
ACCGAUUAAC CAAUGUUCAG CUCACAGUUG CCAGCAAUC A GGGCUAUUU UUUUUUUUU 1500
UUUAUUUUAU GCUAAUUAU UAUAAUUAU UUAGUUUUAU AACUUGCUU UCCUAAAAA 1560
ACCAAUUCGA GAAAGGAACU UUUGCACAGG CAAAAACCU U 1601

SEQ ID NO : 19

SEQUENCE LENGTH : 12

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Cys Phe Thr Ser Ser Tyr Phe Pro Asp Asp Arg Arg

5

10

SEQ ID NO : 20

SEQUENCE LENGTH : 19

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Cys Tyr Thr Ser Ile Glu Lys Tyr Asp Ile Ser Lys Ser Asp Pro

5

10

15

Leu Ala Ala Asp

SEQ ID NO : 21

SEQUENCE LENGTH : 1553

SEQUENCE TYPE : nucleic acid

STRANDEDNESS : double

TOPOLOGY : linear

MOLECULE TYPE : Genomic DNA

SEQUENCE DESCRIPTION :

TTTTACATAT ATTATTCACT CAATATCATA ACAAAAACAA ACTGAATGAT GGCATCTTCT 60

ATTTTGCCTT CCAAATAAT ACAAAAACCG TACCAATTAT TCCACTACTA TTTTCTTCTG 120

GAGAAGGCTC CTGGTTCTAC AGTTACTGAT TTGAATTTG ATACAAACAT ACAAACGAGT 180
 TTACGTAAAT TAAAGCATCA TCATTGGACG GTGGGAGAAA TATTCCATTA TGGGTTTTG 240
 GTTTCCATAC TTTTTTCGT GTTGTGCTT TCAGCAGCTT CATTTCAT AAAATTACCA 300
 ATAATCTTAG CATTGCTAC TTGTTTTTA ATACCCCTAA CATCACAAATT TTTCTTCCT 360
 GCCTTGCCTCG TTTCACTTG GTTGGCATTAA TATTTACGT GTCTAAAT ACCTCAAGAA 420
 TGAAACCAG CTATCACAGT TAAAGTTTA CCAGCTATGG AAACAATTGT GTACGGCGAT 480
 AATTATCAA ATGTTTGGC AACCATCACT ACCGGAGTGT TAGATATATT GGCATGGTTA 540
 CCATATGGGA TTATTCAATT CAGTTCCCA TTTGTACTTG CTGCTATTAT ATTTCATTT 600
 GGGCCACCAG CGGCATTAAG ATCATTGGA TTTGCCTTG GTTATATGAA CTTGCTTGGA 660
 GTCTGATTC AAATGGCATT CCCAGCTGCT CCTCCATGGT ACAAAAACCTT GCACGGATT 720
 GAACCAGCTA ATTATTCAAT GCACGGGTCT CCTGGTGGAC TTGGAAGGAT AGATAAATTG 780
 TTAGGTGTTG ATATGTATAC CACAGGGTT TCCAATTCAAT CAATCATTGTT TGGGGCATT 840
 CCATCGTTAC ATTCAAGGATG TTGTATCATG GAACTGTTAT TTTGTGTTG GTTGTTCGA 900
 CGATTCAAGT TTGTGTGGGT TACATACGCA TCTTGGCTTT GGTGGAGCAC GATGTATTG 960
 ACCCATCACT ACTTTGTCGA TTTGATTGGT GGAGCCATGC TATCTTGAC TGTTTTGAA 1020
 TTCACCAAAT ATAAATATTG GCCAAAAAAC AAAGAAGGCC TTTTCTGTCG TTGGTCATAC 1080
 ACTGAAATTG AAAAATCGA TATCCAAGAG ATTGACCCCTT TATCATAACAA TTATATCCCT 1140
 GTCAACAGCA ATGATAATGA AAGCAGATTG TATACGAGAG TGTACCAAGA GCCTCAGGTT 1200
 AGTCCCCCAC AGAGAGCTGA AACACCTGAA GCATTGAGA TGTCAAATTG TTCTAGGTCT 1260
 AGACAAAGCT CAAAGACTCA GGTTCCATTG AGTAATCTTA CTAACAATGA TCAAGTGCCT 1320
 GGAATTAAACG AAGAGGATGA AGAAGAAGAA GGCGATGAAA TTTCGTCGAG TACTCCTTCG 1380
 GTGTTGAAAG AGCAACCACA GGGTACCAAC TATGCTGCAT CCTCAGCTAC ATCAGTAGAT 1440
 GATTTGGATT CCAAAAGAAA TTAGTAAAC ACCAGTTCT ATTAAATTCT TTATTCCTC 1500
 CTAATTAAATG ATTTATGTT CAATACCTAC ACTATCTGTT TTTAATTCC TAC 1553

SEQ ID NO : 22

SEQUENCE LENGTH : 472

SEQUENCE TYPE : amino acid

STRANDEDNESS : single

TOPOLOGY : linear

MOLECULE TYPE : peptide

SEQUENCE DESCRIPTION :

Met Met Ala Ser Ser Ile Leu Arg Ser Lys Ile Ile Gln Lys Pro
1 5 10 15
Tyr Gin Leu Phe His Tyr Tyr Phe Leu Leu Glu Lys Ala Pro Gly
20 25 30
Ser Thr Val Ser Asp Leu Asn Phe Asp Thr Asn Ile Gln Thr Ser
35 40 45
Leu Arg Lys Leu Lys His His His Trp Thr Val Gly Glu Ile Phe
50 55 60
His Tyr Gly Phe Leu Val Ser Ile Leu Phe Phe Val Phe Val Val
65 70 75
Phe Pro Ala Ser Phe Phe Ile Lys Leu Pro Ile Ile Leu Ala Phe
80 85 90
Ala Thr Cys Phe Leu Ile Pro Leu Thr Ser Gln Phe Phe Leu Pro
95 100 105
Ala Leu Pro Val Phe Thr Trp Leu Ala Leu Tyr Phe Thr Cys Ala
110 115 120
Lys Ile Pro Gin Glu Trp Lys Pro Ala Ile Thr Val Lys Val Leu
125 130 135
Pro Ala Met Glu Thr Ile Leu Tyr Gly Asp Asn Leu Ser Asn Val
140 145 150
Leu Ala Thr Ile Thr Thr Gly Val Leu Asp Ile Leu Ala Trp Leu
155 160 165
Pro Tyr Gly Ile Ile His Phe Ser Phe Pro Phe Val Leu Ala Ala
170 175 180
Ile Ile Phe Leu Phe Gly Pro Pro Thr Ala Leu Arg Ser Phe Gly
185 190 195

Phe Ala Phe Gly Tyr Met Asn Leu Leu Gly Val Leu Ile Gin Met
200 205 210
Ala Phe Pro Ala Ala Pro Pro Trp Tyr Lys Asn Leu His Gly Leu
215 220 225
Glu Pro Ala Asn Tyr Ser Met His Gly Ser Pro Gly Gly Leu Gly
230 235 240
Arg Ile Asp Lys Leu Leu Gly Val Asp Met Tyr Thr Thr Gly Phe
245 250 255
Ser Asn Ser Ser Ile Ile Phe Gly Ala Phe Pro Ser Leu His Ser
260 265 270
Gly Cys Cys Ile Met Glu Val Leu Phe Leu Cys Trp Leu Phe Pro
275 280 285
Arg Phe Lys Phe Val Trp Val Thr Tyr Ala Ser Trp Leu Trp Trp
290 295 300
Ser Thr Met Tyr Leu Thr His His Tyr Phe Val Asp Leu Ile Gly
305 310 315
Gly Ala Met Leu Ser Leu Thr Val Phe Glu Phe Thr Lys Tyr Lys
320 325 330
Tyr Leu Pro Lys Asn Lys Glu Gly Leu Phe Cys Arg Trp Ser Tyr
335 340 345
Thr Glu Ile Glu Lys Ile Asp Ile Gln Glu Ile Asp Pro Leu Ser
350 355 360
Tyr Asn Tyr Ile Pro Val Asn Ser Asn Asp Asn Glu Ser Arg Leu
365 370 375
Tyr Thr Arg Val Tyr Gln Glu Pro Gln Val Ser Pro Pro Gln Arg
380 385 390
Ala Glu Thr Pro Glu Ala Phe Glu Met Ser Asn Phe Ser Arg Ser
395 400 405
Arg Gln Ser Ser Lys Thr Gln Val Pro Leu Ser Asn Leu Thr Asn

410 415 420
Asn Asp Gln Val Pro Gly Ile Asn Glu Glu Asp Glu Glu Glu
425 430 435
Gly Asp Glu Ile Ser Ser Ser Thr Pro Ser Val Phe Glu Asp Glu
440 445 450
Pro Gln Gly Ser Thr Tyr Ala Ala Ser Ser Ala Thr Ser Val Asp
455 460 465
Asp Leu Asp Ser Lys Arg Asn
470

[Claims]

[Claim 1] An isolated gene coding for a protein which regulates aureobasidin sensitivity.

[Claim 2] An isolated gene as claimed in Claim 1 which is contained in a DNA fragment represented by a restriction enzyme map as specified in any of Fig. 1 to Fig. 3.

[Claim 3] An isolated gene as claimed in Claim 1 which is hybridizable with a gene of Claim 2.

[Claim 4] A process for cloning a gene of Claim 1 characterized by using a gene of Claim 2 or 3 or a part thereof as a probe.

[Claim 5] A nucleic acid probe which comprises a sequence consisting of 15 or more bases and is hybridizable with a gene of Claim 1.

[Claim 6] An antisense DNA of a gene which codes for a protein regulating aureobasidin sensitivity.

[Claim 7] An antisense RNA of a gene which codes for a protein regulating aureobasidin sensitivity.

[Claim 8] A recombinant plasmid containing a gene of Claim 1.

[Claim 9] A transformant having a recombinant plasmid of Claim 8 introduced thereinto.

[Claim 10] A process for producing a protein regulating aureobasidin sensitivity characterized by culturing a transformant of Claim 9 and collecting the protein regulating aureobasidin sensitivity from the culture.

[Claim 11] An isolated protein regulating aureobasidin sensitivity which is encoded by a gene of Claim 1.

[Claim 12] An antibody against a protein of Claim 11.

[Claim 13] A process for detecting a protein regulating aureobasidin sensitivity which comprises using an antibody of Claim 12.

[Claim 14] A process for detecting a gene coding for a protein regulating aureobasidin sensitivity which comprises the hybridization with the use of a nucleic acid probe of Claim 5.

[Claim 15] A process for screening an antimycotic which comprises using a transformant of Claim 9 or a protein of Claim 11.

[Claim 16] An isolated gene coding for a protein which regulates aureobasidin sensitivity substantially as herein described with reference to any one of the Examples.

[Claim 17] A transformant having an isolated gene coding for a protein which regulates aureobasidin sensitivity substantially as herein described with reference to any one of the Examples.

[Claim 18] A process for producing a protein which regulates aureobasidin sensitivity substantially as herein described with reference to any one of the Examples.

Dated this 16th day of May 1994

TAKARA SHUZO CO., LTD.
By their Patent Attorney
GRIFFITH HACK & CO.

[Designation of Document] Abstract

[Abstract]

[Object] To provide a protein regulating the sensitivity to an antimycotic aureobasidin, a gene coding for this protein, the use thereof, an antibody for the protein and the use thereof.

[Constitution] An isolated gene coding for a protein regulating aureobasidin sensitivity. A process for cloning the gene with the use of the gene or a part of the same as a probe. A nucleic acid probe being hybridizable with the gene. An antisense DNA or RNA of the gene. A recombinant or transformant having the gene contained therein. An isolated protein regulating aureobasidin sensitivity and a process for producing the same by using the transformant. An antibody for the protein. A process for detecting the protein or the gene. A process for screening an antimycotic by using the protein or the transformant.

[Effects] Useful in the diagnosis and treatment for diseases including mycoses.

[Selected Figure] none.

Fig. 1

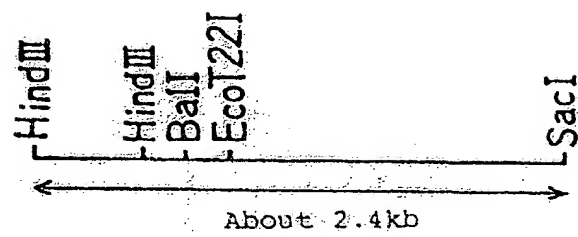


Fig. 2

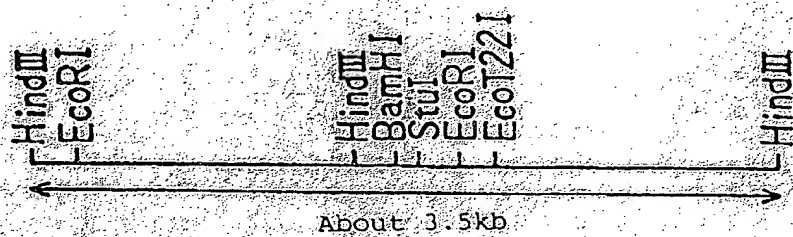


Fig. 3

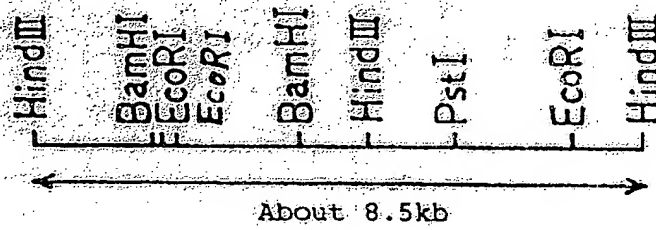


Fig. 4

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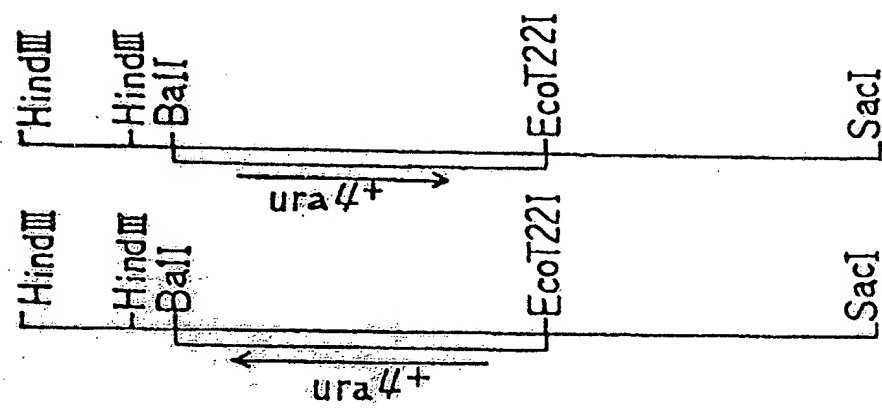
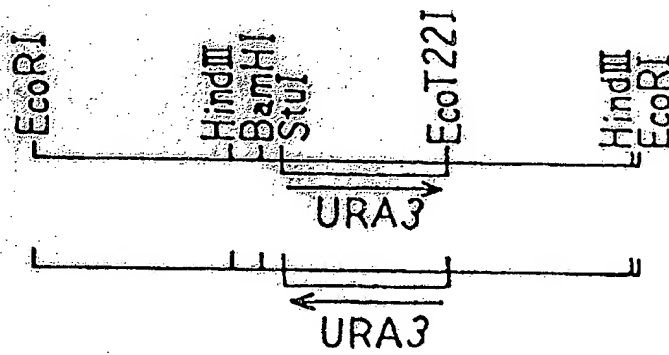


Fig. 5



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Fig. 6

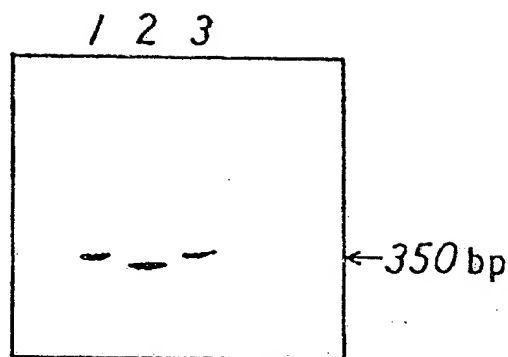


Fig. 7

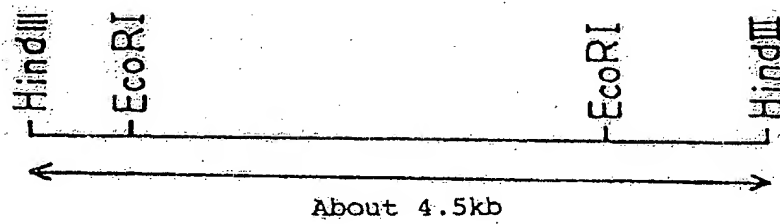


Fig. 8

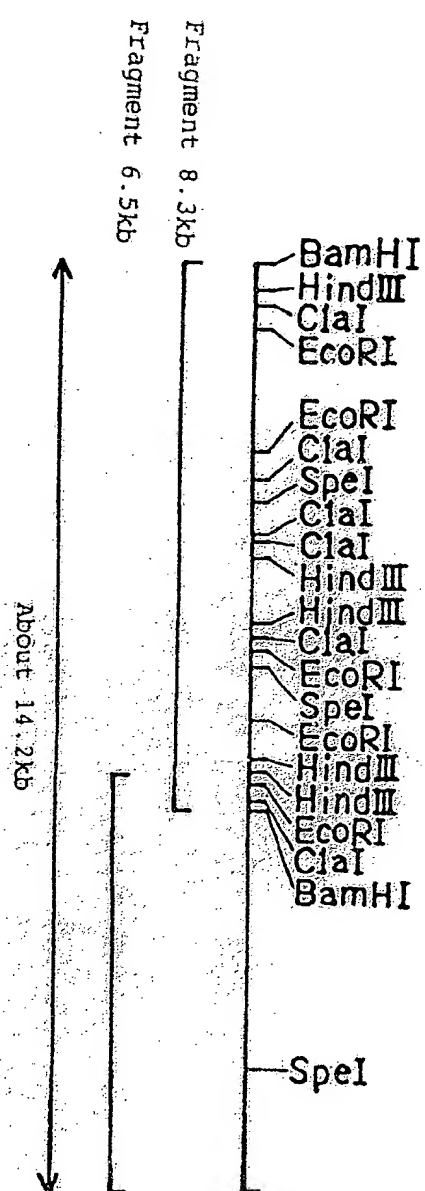
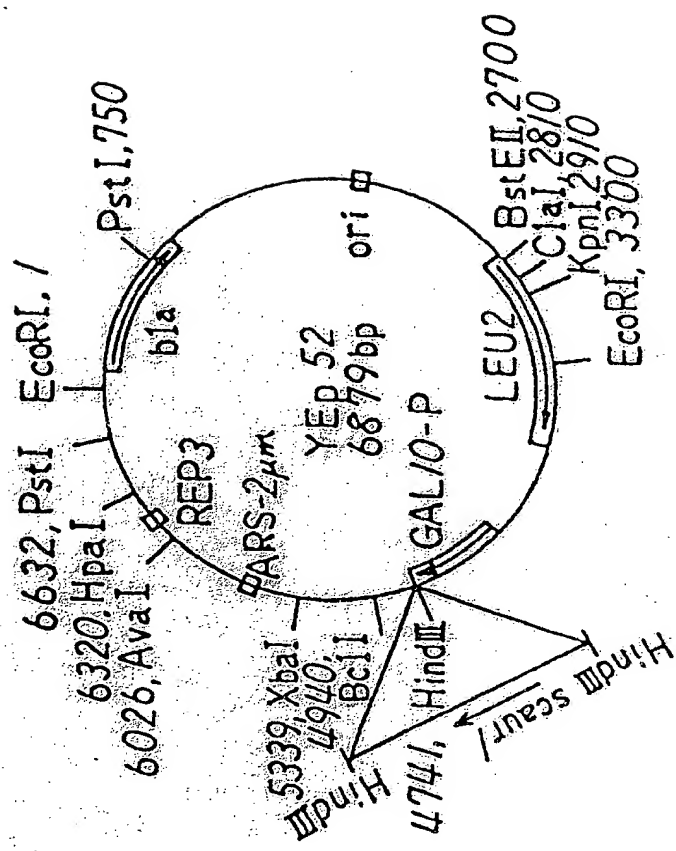


Fig. 9



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Fig. 10

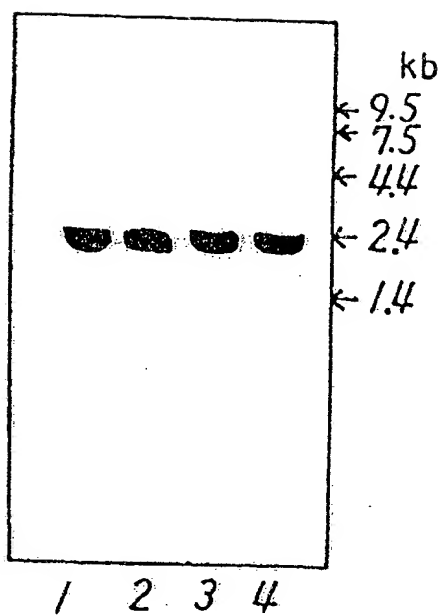


Fig. 11

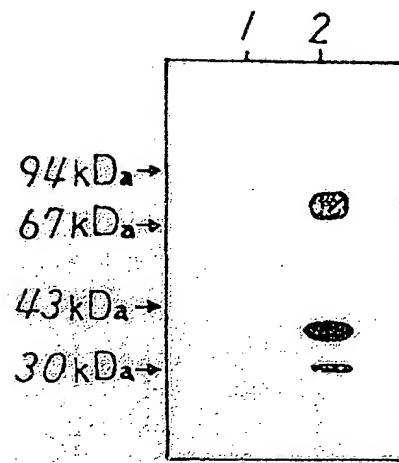


Fig. 12

